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Committee on Health, Education, Labor, and Pensions

United States Senate

One Hundred Sixth Congress

First Session

On

Vermont Field Hearing on Federal Legislation Relating to Medical Information Confidentiality

March 15, 1999 (Berlin, VT.)

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(III)
MEDICAL RECORDS PRIVACY

MONDAY, MARCH 15, 1999

U.S. SENATE,
COMMITTEE ON HEALTH, EDUCATION, LABOR, AND PENSIONS,
Berlin, VT.

The committee met, pursuant to notice, at 9:19 a.m., in Conference Room 3, Central Vermont Medical Center, Berlin, Vermont, Senator James M. Jeffords (chairman of the committee) presiding. Present: Senator Jeffords.

OPENING STATEMENT OF SENATOR JEFFORDS

The CHAIRMAN. Good morning. May I have your attention please. As you know, I am Senator Jim Jeffords. I am chairman of the Health, Education, Labor, and Pension Committee, called the HELP Committee, and right now it is the busiest legislative committee in the Senate. And we are here today to hold a hearing, which I try to do in Vermont as often as I can, especially if the subjects are of great importance to individuals, to businesses, and to hospitals.

Senator Leahy should be joining us before too long. Senator Leahy and I have been in the Congress about 24 years together now and have learned to work together very well. He is on the Judiciary Committee, which also has an interest in the legislation which we are talking about today.

This is the committee's sixth hearing on the topic of medical records privacy. And until recently we gave very little thought to issues of medical privacy. We visited our family doctor with full confidence in the sanctity of the physician-patient relationship. Today, however, technology has radically altered the delivery of health care and the use of health care information. The National Research Council, as part of their study of the confidentiality of medical records, has shown that numerous individuals in more than 17 different organizations have access to any one set of medical records.

Many recent technology changes contribute to better and more efficient health care. Telemedicine will make it possible for patients here at Central Vermont Hospital to benefit from the expertise of physicians at nationally known academic medical centers like Fletcher Allen, Dartmouth-Hitchcock.

Quality care requires more than the free flow of information between providers, payers, and other users of health information. It requires trust between a patient and a caregiver. For our health care system to be effective as well as efficient, patients must feel comfortable sharing sensitive information with health profes-
sionals. Technology has provided the tools to allow the ease of access to health care information. Now legislation is needed to ensure the confidentiality of personal health information.

The current loose web of State privacy laws is inadequate and insufficient. We need a uniform national standard for medical records confidentiality, with the exception of State laws relating to public health issues in those areas having a history of discrimination, such as mental health. The Health Insurance Portability and Accountability Act of 1996, known as the Kassebaum-Kennedy Act, established several mandates relating to medical records privacy. One provision set August, 1999, this August, as the deadline by which Congress must act to ensure the confidentiality of electronically submitted data. If for some reason Congress fails to act, then HIPAA includes a default provision directing the Secretary of Health and Human Services to promulgate regulations.

This issue is much too complex and important to be handled by the regulatory process. Senator Leahy and I prefer to ensure the appropriate safeguards for medical data through the deliberative and conscientious approach of the legislative process. This hearing is intended to be an overview of the unique perspectives of consumers, providers, regulators, insurers, employers, and researchers. I look forward to all the valuable testimony today that we have before us so that we can move forward to meet that August deadline.

I would like to announce that following the three panels of witnesses, we will have an open mike session for questions and comments. There is a sign-up sheet for use of the microphone in the back of the room. Since we have a limited amount of time, sign-up will be on a first-come-first-served basis, and we will listen to as many people as we possibly can.

We will be hearing from several distinguished witnesses this morning. On the first panel, I am happy to introduce Thomas Koch, who has been elected to his fourth term as the Republican member of the Vermont House representing Barre Town. Is Mr. Koch here? Oh, yes, I see you now. Hi. How are you?

An attorney, Tom Koch also has served since 1984 as Barre Town moderator. If you want to come up, Tom, and take a seat. He was a member of the Barre Town Charter Committee. Tom has been active in a wide variety of community service roles, having served with the Army in Vietnam, with the Barre Lions Club, and as a scoutmaster. It is a great pleasure to have you with us this morning, and thank you for appearing.

The next is Ms. Meg O'Donnell, director of quality assurance and consumer protection for the Vermont Division of Health Care Administration. In that capacity she holds responsibility for implementing Rule 10, Vermont's new regulatory oversight program for managed care plans. In the course of her career she has frequently spoken on health care policy topics such as consumer protections and the confidentiality of medical records. Ms. O'Donnell, we are pleased to have the benefit of your experience.

The next witness, whom I am pleased to introduce is Susan L. Aranoff, a staff attorney at Vermont Protection and Advocacy, Inc., in Montpelier, where she provides legal services to people with disabilities and also is engaged in policy research, development, and advocacy before the legislature and other agencies. In addition, Ms.
Aranoff is co-chair of the Vermont Coalition for Disability Rights, Legislative Committee.

In the past she served clerkships in the office of Governor Kunin and the office of Vermont's Attorney General.

Ms. Aranoff, we thank you for being with us today.

Also testifying today is Mr. Tim Palmer, executive director of Vermont Cares in Burlington. Vermont Cares is a statewide organization committed to providing resources, education, and services for people with HIV and AIDS.

Mr. Palmer, thank you for being here. Good to see you again.

Tom, would you please proceed.

STATEMENTS OF HON. THOMAS KOCH, CHAIRMAN, WAYS AND MEANS COMMITTEE, VERMONT STATE HOUSE OF REPRESENTATIVES, BARRE TOWN, VT; MEG O'DONNELL, DIRECTOR OF QUALITY ASSURANCE AND CONSUMER PROTECTION, VERMONT DIVISION OF HEALTH CARE ADMINISTRATION, MONTPELIER, VT; TIM PALMER, EXECUTIVE DIRECTOR, VERMONT CARES, BURLINGTON, VT; AND SUSAN ARANOFF, ATTORNEY, VERMONT PROTECTION AND ADVOCACY, MONTPELIER, VT

Mr. Koch. Thank you, Mr. Chairman. I thank you for the opportunity to share some thoughts with you on the subject of medical records confidentiality. This is not a high profile issue, and few people seem concerned with it until the relatively rare event occurs which brings it home personally. The fact is, however, that everyone has some form of medical record, and everyone is interested in maintaining some degree of control over the dissemination of his or her own records. So I thank you for your interest in this matter and the important work that you are doing on it.

This issue, like so many that Congress on the one hand and State legislatures on the other deal with, presents a difficult question of Federal-State relationships. I believe I speak for 179 other members of the Vermont General Assembly when I tell you that we are jealous of our legislative prerogatives and that we believe we are in a better position than the Congress to tailor programs and regulations in a manner which best serve and protect Vermont's citizens. On the other hand, we certainly recognize the constitutional role of Congress to regulate commerce among the States, and that is an area that has become vast more complex in recent years. The proper balance point is not easily located, and I do not presume to find it for you today. I will, however, attempt to share what we are doing in Vermont and to express some concerns, and I hope this will be helpful to you as you work on the subject in Washington.

To the extent that Federal law will preempt State law, it would be useful for you to know what it is that you are preempting, and I wish I could tell you. Vermont's medical records confidentiality laws are not a coherent whole. Rather, they have grown up piecemeal over many years, and rather than being comprehensive, they could best be described as hit and miss. We do have statutes and court rules relative to provider-patient privilege, but there is some question of the extent to which records, as opposed to direct communication, are privileged. The privilege has been substantially
narrowed by a number of court decisions over the years. Requirements regarding the length of time that records must be maintained, except hospital records, do not exist. And Vermont's laws certainly do not contemplate the ease with which information can be transferred in the age of the Internet.

So the need for a new medical records confidentiality law in Vermont has been clear for a number of years. Nevertheless, the bill has foundered on the rock of complexity during the past two biennial sessions, and we are just now beginning the process again. Last week, the House Health and Welfare Committee, on which I serve, voted out its version of the bill, H. 63, and that bill is now in the House Judiciary Committee. Eventually, I expect the corresponding committees in the Vermont Senate to consider the bill as well. For that reason, I cannot tell precisely what the final version will contain.

What I can tell you is a lot of work has gone into this bill with more to come. We have listened to many, many Vermonters trying to design a law that will serve their particular needs. We have attempted to design a bill that is both comprehensive and workable. How any other State may view the needs of their people and what other states have done in this regard, I do not know. We have not, however, heard any concerns in our committee expressed to us that what another State has done might adversely impact Vermonters. For that reason, I do not perceive any need for Congress to roll back, by way of preemption, any protections that Vermont or any other State has adopted.

There are those who may argue that some businesses, such as insurance companies who deal with medical records across State boundaries, and reviewing some of the other statements that will be made here today, physicians, for example, who treat people across State boundaries, will find it difficult to comply with a myriad of regulations imposed by 50 different states. I would suggest that it will not be any more difficult than it is for insurers to comply with existing insurance regulations in the several states. They have developed State specific endorsements, procedures and protocols in the conduct of their business, and they will no doubt do so as they deal with modern medical records confidentiality laws. In short, this is a matter in which Congress may be tempted to impose uniformity, but I would hope the temptation would be resisted.

What it does raise is I believe a legitimate concern are the truly Federal questions about transfer of information across State lines, particularly to a person in a State which may not have acted adequately to maintain the confidentiality of records in its jurisdiction. Vermont's proposed bill, for example, makes it clear that a person who or which obtains health care information for a lawful purpose is a custodian of that information, and subsequent disclosure of that information is subject to the provisions of the proposed act. When the recipient is in another State, it is unclear how Vermont can enforce the protections it purports to give to its citizens. In this case, the Federal power can be helpful, either by enacting minimum standards where inadequate rules exist, or by giving effect in Federal law to the law of the jurisdiction in which the medical in-
formation originated if that state’s protections are greater than those of the State to which the information is sent.

The Federal-State question is sometimes reduced to the shorthand whether the Federal law should be a floor or a ceiling. My own conclusion is that there is no pressing need for uniformity, that the States should be allowed to respond to the need of the citizens as the 50 State legislatures deem best, and that if a State provides greater protections to its citizens than are provided under whatever law the Congress eventually enacts, the additional protection ought to be—ought to remain in effect, thus, providing a Federal floor but leaving the states to establish the ceiling.

I thank you for this opportunity and for your consideration.

The CHAIRMAN. Thank you very much. I think we will go down through the statements and then I will have questions.

The CHAIRMAN. Ms. O’Donnell.

Ms. O’DONNELL. Thank you. Good morning, Mr. Chairman, and thank you for this opportunity to address your committee on the confidentiality of medical information.

My name is Meg O’Donnell, and I am currently the director of quality assurance and consumer protection for the division of health care administration which is part of the Vermont Department of Banking, Insurance, Securities and Health Care Administration. In this and my previous position as general counsel to the Health Care Authority, I have been interested and involved for a number of years in the issue of ensuring the confidential handling of individually identifiable health care information.

First, I want to say that we are very proud that Vermont’s two senators have been so involved in this issue and addressing this critical issue on the national level. We are pleased that the work continues and very pleased that we have this opportunity to address you at today’s hearing.

Vermont is not unique in its current approach to medical records confidentiality. As Representative Koch I think amply illustrated, we have a hodgepodge of laws that provide some protections, but certainly not the comprehensive protections that Vermonters can and should expect in the area of their medical records.

The division has for some time now recognized a need for broad legislation that will comprehensively govern how medical information is collected, used and maintained. Our involvement in this issue is based in part on the division’s own need for clinical data to carry out some of our legally mandated activities. For example, the division contracts with the Vermont Program for Quality in Health Care to develop Vermont specific practice guidelines which requires an understanding of how health care has been delivered, and that depends on medical records. Another example is the division’s oversight of managed care plans. By law, we are required to audit such plans at least once every 3 years. Those audits must include a review of a selected number of medical records in order to ensure that plan practices comply with our regulations.

Vermonters must have confidence that the Division of Health Care Administration and other public entities engaged in similar work are handling their records with the respect, security and confidentiality they expect. While I believe that most health care providers, facilities and insurers in Vermont are already setting and
abiding by high standards in handling their medical information, the public’s confidence can only be increased by setting a benchmark standard that all people and organizations who have access to medical records must legally comply with.

To that end, the division has supported the efforts of the Vermont legislature over the past 4 years to pass a comprehensive law setting basic standards for how confidential medical records are used.

But even if Vermont succeeds in passing one of the bills presently before it, a State-by-State approach to medical records confidentiality will not solve all of the issues. Health care is no longer a matter that fits neatly within State borders. Again, Vermont is an excellent example of this. 40 percent of the patients seen at Dartmouth-Hitchcock Medical Center actually live in Vermont. Similarly, many Vermonters have sought or will at some time seek care even further afield at centers of excellence located in Boston, in New York, in Cleveland, in any number of other states. The reality of the health delivery system today is that it is a multistate system. So no matter what one State, like Vermont, accomplishes in protecting its residents’ medical records, those protections will not stretch as far as those medical records are likely to travel.

Thus, while we continue to support Vermont legislation in this area, we strongly support action on the part of the Congress on this critical issue. Only Federal legislation can ensure that all confidential medical records receive the same basic protections no matter where in the United States they are located, used or transmitted.

Any such legislation must strike a balance between privacy rights of individuals and the legitimate use of individually identifiable information in delivering and paying for health care services in research and in overseeing the quality of today’s health care system. Essential ingredients of such legislation include, and I will just list them very briefly, patients’ rights. We believe that individuals should have primary control over disclosure of their health care information. Standards for releasing identifiable health care information. Standards for maintaining identifiable health care information. And as importantly, if not more importantly, effective sanctions.

Let me reiterate a concept that the division considers essential to any proposed legislation, and I apologize, I have not had a chance to read S. 578 so I cannot speak in any detail about that.

The CHAIRMAN. I understand, Ms. O’Donnell.

Ms. O’DONNELL. In the desire to protect individuals’ medical records from unnecessary release or use, we must ensure that legitimate uses can continue without undue burden. From a regulator’s perspective, for example, this includes the ability to review medical records as necessary to carry out my legal responsibilities. A good law will allow this use, but will also establish strong sanctions for any misuse or abuse that could result.

We look forward to the opportunity to review and comment in more detail on S. 578, and I thank you again.

[The prepared statement of Ms. O’Donnell follows:]
Thank you for this opportunity to address the Committee on the confidentiality of medical information.

My name is Meg O’Donnell, and I am currently the Director of Quality Assurance and Consumer Protection for the Division of Health Care Administration, part of the Vermont Department of Banking, Insurance, Securities and Health Care Administration. I was formerly the General Counsel to the Health Care Authority, which became the Division of Health Care Administration in 1996 as the state consolidated its regulatory oversight of health delivery systems and insurance companies in Vermont. In both positions, I have been interested and involved for a number of years in the issue of ensuring the confidential handling of individually-identifiable health care information.

First, I want to say how proud I am that Vermont’s two senators have been at the forefront of addressing this critical issue on the national level. I am pleased that this work continues to be a focus of their offices and this Committee, as evidenced by today’s hearing.

Vermont is not unique in its current approach to medical records confidentiality. While many, if not most, consumers believe that their medical records are uniformly subject to tight legal restrictions on how they can be used, Vermont in fact has a medley of laws that address only bits and pieces of the issues. For example, we have a Hospital Bill of Rights that protects medical records of inpatients, and our managed care oversight program has regulatory provisions that govern how managed care organizations handle their patients’ confidential records. But these and similar provisions do not provide the comprehensive protections Vermonters can, and should, expect.

The Division has for some time now recognized a need for broad legislation that will comprehensively govern how medical information is collected, used and maintained. Our involvement in this issue is based on the Division’s own need for clinical data to carry out some of its legally mandated activities. For example, the Division contracts with the Vermont Program for Quality in Health Care to develop Vermont specific practice guidelines, which requires an understanding of how care has been delivered in the past—and that depends on medical records. Another example is the Division’s oversight of managed care plans. By law, we are required to audit such plans at least once every three years. Those audits must include a review of a selected number of medical records in order to ensure that plan practices comply with our regulations.

Vermonters must have confidence that the Division of Health Care Administration and other public entities engaged in similar work are handling their records with the respect, security and confidentiality they expect. While I believe that most health care providers, facilities and insurers in Vermont already set and abide by high standards in handling medical information the public’s confidence can only be increased by setting a benchmark standard that all people and organizations who have access to medical records must legally comply with.

To that end, the Division has supported the efforts of the Vermont legislature over the past four years to pass a comprehensive law setting basic standards, for how confidential medical records are used.

But even if Vermont succeeds in passing one of the bills presently pending before it, a state-by-state approach to medical records confidentiality will not solve all of the issues. Health care is no longer a matter that fits neatly within state borders. Vermont is an excellent example of this, forty percent of the patients seen at the Dartmouth-Hitchcock Medical Center, which is located across the border in New Hampshire, actually live in Vermont. Similarly, many Vermonters have sought or will at some time seek care even further afield, at centers of excellence located in Boston, in New York-, in Cleveland, in any number of other states. The reality of the health delivery system today is that it is a multistate system. So no matter what one state, like Vermont, accomplishes in protecting its residents’ medical records, those protections will not stretch as far as those medical records are likely to travel.

Thus, while we continue to support Vermont legislation in this area, we strongly support action on the part of the Congress on this critical issue. Only federal legislation can ensure that all confidential medical records receive the same basic protections no matter where in the United States they are located, used or transmitted.

Any such legislation must strike a balance between the privacy rights of individuals and the legitimate use of identifiable information in delivering and paying for health care services and oversee in the quality of today’s health care system. Essential ingredients of such legislation include:
• **Patients’ rights.** Individuals should have primary control over disclosure of their health care information. Those rights should include the right to consent to disclosures before they are made, except in certain defined circumstances specifically authorized by law, the right of patients to see their own health care records and to receive copies of them without being charged excessively high fees, and the right to correct health-care records that contain wrong information.

• **Standards for releasing identifiable health care information.** Those standards should clarify and codify the circumstances under which health care providers and facilities may release health care information without the individual’s consent, and establish a general prohibition against disclosing or re-disclosing identifiable health care information without a consent or specific legal authority.

• **Standards for maintaining identifiable health care information.** The law should establish basic standards for maintaining the security and confidentiality of health care records, whether by a health care facility, provider or insurer.

• **Effective sanctions.** Legislation must include effective enforcement tools.

Let me reiterate a concept that the Division considers essential to any proposed legislation. In the desire to protect individuals’ medical records from unnecessary release or use, we must ensure that legitimate uses can continue without undue burden. From a regulator’s perspective, for example, this includes the ability to review medical records as necessary to carry out my legal responsibilities. A good law will allow this use, but will also establish strong sanctions for any misuse or abuse that could result.

We look forward to the opportunity to review and comment in more detail on any proposed legislation that the Committee will be considering.

Thank you.

The CHAIRMAN. Thank you very much. It is very helpful.

Mr. Palmer.

Mr. PALMER. Thank you, Senator. Thank you for this opportunity to address this important issue. I would like to indicate for the record that the airlines ate my testimony. My testimony is in a suitcase in Philadelphia. [Laughter.]

The CHAIRMAN. That may be helpful there. [Laughter.]

Ms. ARANOFF. Air passenger's Bill of Rights theory?

Mr. PALMER. Air passenger’s Bill of Rights. I will be submitting my written remarks shortly, but first of all, I wanted to address the fundamental issue of why I am testifying; that is, that for the past 2 weeks we have tried to identify someone living with HIV in the State of Vermont who will be willing to testify on this subject. Given the public nature of this hearing, all of the people that we work with on a regular basis have declined this opportunity. That is how important confidentiality is to people living with HIV.

So I am put in the position of reporting that on a regular basis people living with HIV are faced with making health care decisions based on what the record is ultimately going to show; who is going to get that record. That is a barrier to health care. And I believe that your legislation moves in the right direction in providing, as Representative Koch has indicated, a Federal floor. And I would encourage you—in support of Representative Koch’s position, I would encourage Federal legislation to be the floor and to permit states to build over, above and beyond that floor.

I think it is important to recognize that people live with life threatening and chronic illness and with those illnesses also including a great deal of judgment about their lives and their own personal judgments, it is important that—that their health care be something that is above and beyond the probing of outside experts.

It is important to recognize that that barrier to care has prevented and adds an additional burden for someone who may be the victim of a sexual assault from accessing the kinds of care that are in that survivor’s best interest because of fear that at some future
point an employer accessing those records can find out that that individual sought testing around HIV and was treated. Not because of any action taken by that individual but because it was in their best interests to protect their health and safety. That is just one example of how important it is to maintain the confidentiality of records, particularly when it comes to employment situations.

It is important to establish in the law that employers need to make decisions on employment as people living with HIV go back to work, as your—as another bill that you have advanced addresses. As they go back to work, it is important that employers make decisions based on the individual's qualifications and ability, not on their medical past. And I encourage that that kind of issue be addressed in your legislation. And I thank you for the opportunity to address you today.

The CHAIRMAN. Thank you, Ms. Aranoff.

Ms. ARANOFF. Good morning, Senator. I also would like to thank you and acknowledge the hard work that you have done over the years to bring this issue of medical records and the need for confidentiality to the attention of Congress, and the work that you did last year to introduce a bill, and I understand you recently reintroduced a new bill, that has, I think, really moved this issue forward. And I know that you have galvanized a lot of this attention that you have received at the Statehouse here in Montpelier as well. So I want to thank you for all your work on that issue.

Before I begin my testimony, on behalf of Vermont Protection Advocacy I want to share with you a story that was my first introduction to this issue of medical records confidentiality. And we were asked to try to find a person whose life has been affected either by not being able to get access to their own medical record or by a breach of confidentiality of their own medical record. And in 1992 when I was an attorney in private practice, by just a fluke of nature the Vermont Lawyer Referral Service sent my office a case that involved a woman, she is now deceased. And this is all a matter of public record, and I think if she were here, she would either be sitting here today or saying, yeah, go ahead and tell my story because it was a really compelling story and it illustrates the need for this exact legislation.

This woman was a Vermonter, and she had one of those childhoods that we read about. But you actually meet someone who has experienced sexual abuse by relatives, placement in foster care, sexual abuse by foster parents, ending up at Brattleboro Retreat for many drug overdoses, alcohol issues by the time she was 19, and then being released from Brattleboro Retreat to an interim, like halfway facility called Spruce Mountain Inn here in this area, receiving some treatment at Spruce Mountain Inn and then being discharged presumably into the community which she ended up homeless—at the point when I met her she was 22, and she was living next door to her mother in a duplex in Plattsburgh, New York. She was taking care of her mother who had lung cancer and was in the very end stages of a long and painful illness.

And in the course of taking care of her mother she was doing routine things like cleaning the house, vacuuming and stuff like that. She was vacuuming under her mother's bed 1 day and saw a stack of papers and went to move them and realized that they
were her treatment records from both the Brattleboro Retreat and Spruce Mountain Inn. And she panicked because at facilities—at mental health facilities a lot of times what goes into a treatment record are the person's own handwritten statements. This was around the time when David Letterman came out with his top ten lists. So she had a lot of top ten lists that were assignments. The top ten things I am most ashamed of, the top ten things I have never told anyone, the top ten reasons I am mad at my mother. Her mother was dying of lung cancer caused by cigarette smoking, and this woman was really pissed that her mother smoked her whole life and was going to be dead by the time she was 25. And that was like her top ten reason for being mad at her mother.

In the top ten things that she had never told anyone and was most ashamed of was a lot of this history of family sexual abuse and other kinds of abuse that she had never told her mother. And so there she is cleaning and there she discovers that all of this very sensitive, very—well, I will leave it at sensitive information that she had provided to treatment providers that she had trusted had been released to her mother.

Her mother died within 6 months of that, and during that time they tried to deal with some of these issues, but it kind of complicated her grieving process with her mother. She came to my office wanting to know, could she sue over this. What—did she have any recourse? In the course of doing this litigation, we ran into issues like definition of medical records. None of this information that she provided—none of this information that was released, the stuff that really caused her harm, was written by a provider. It was not a provider's treatment notes, it was her own statements, and it was not necessarily about a medical condition. So the definitions that we have in the bill that is under consideration in Vermont and in DC. would not necessarily capture this information that was created in the course of receiving treatment and stored and maintained by a treatment provider and released by the treatment provider, would not necessarily be within the definition of medical information.

Perhaps, you know, significant to an attorney, but I think to others as well, is the definition of harm and the fact that when there is a release of this type, there is very little recourse right now. We need like significant penalties for negligent releases as well as intentional releases. But because she was a person living on disability, and because she did not have lost wages, and because it is hard to quantify the injury to a mother-daughter relationship, you know, the 6 months—the last 6 months of that person's life, she did not have really good damages, you know, from like a litigator point of view. And so—and there was no other penalty to kind of capture that case.

So anyway, that was a lesson of that litigation. The need for remedies, the need for definitions, the need for—the other thing we learned about both of those facilities was they had no policies or procedures in place for how they treat the journals, and diaries and assignments, writing assignments that they give their patients. They did not have policies as to whether or not they considered that medical information, whether or not they released that. The patient kept it in a file, anything like that. And how long they
would keep it for, and what they would do—what they would send it to when they could not find anyone. In this case she was homeless, they did not have a known address. They sent it to her mother.

A lot of people would think what is the big deal, but when it happens to someone, it can be a really big deal. Her mother had no knowledge. So since I have used up most of my time telling that story, I will hit the high points of my testimony.

The CHAIRMAN. Could you speak a little louder? Imagine you are talking to the back of the room instead of to me. I understand that is hard. There are a number of people here who would like to hear from all of you.

Ms. ARANOFF. Our agency is federally funded to assist people just like the person I described, people with diagnoses, labels, history of mental illness who are receiving residential care. We keep statistics on the number of—on the issues that come up and the cases that we handle as well as on the issues in the cases that we do not handle but that we provide information, referrals and services to.

In 1997, 1998 and to date in 1999 we have had a total of 28 clients with—these are clients, not information referrals, these are people we have actually opened up cases for who have had issues around their medical records. Fifteen of those have been breaches of confidentiality and 13 of those have been access—access issues. And the access issue is probably the overarching issue for our clients. Very often they are denied access entirely. Sometimes their access is restricted.

And I brought with me a copy of this letter that is dated February 24th, 1999, so this is recent. This is what happens today when someone asks for their treatment notes. It is from Counseling Service of Addison County, and the response to a person’s request for their record is to say, first, we need to meet with you to go over the record together before you can copy anything to take. And I will be glad to schedule a meeting with you, but in anticipation of this meeting, please let me know the reason for your wanting these records.

Now these kinds of hurdles might get placed in the way of people with other kinds of conditions other than mental illness, but they are constantly placed in the way of people who want their mental health treatment records. Sometimes it is an outright denial for that category, and the bill that was first introduced in the Vermont legislature would have sanctioned a denial of records access for people with mental illness. Access issues are key.

Along with the outright denials of records, there are other kinds of obstacles, and one of those is financial. Brattleboro Retreat charges $1 a page for copying. We have a client whose record from a 30-day stay is estimated to cost more than $1,500 just to get a copy of a record, and that is not an expense that insurance will reimburse. The State Hospital, which is supposed to under Vermont law charge only 4 cents a page, routinely charges 50 cents a page. When you are there for 30 days, you know, 60 days, that kind of cost can become quite prohibitive. We think that everyone should be entitled to one free copy of their record, and that if their record
is needed for a benefits case or something like that, that it should be free as well.
In addition to access issues, the other complaint—the other situation that affects our clients most is confidentiality. One of the most common things—it shocks me all the time that I can call up on the phone and talk to insurers and treatment providers about someone's mental health treatment and I am not even asked if I have a release, let alone asked to potentially fax the release. This happens to me in my office I would say easily five times a week. And I think that those kinds of things happen.
I have a written statement that has addressed these issues in more detail and, again, I thank you for your time and attention to this issue.

The CHAIRMAN. Thank you for an excellent statement. Let me welcome Senator Leahy. We have worked together for about 24 years now in Congress and the Senate, and it is a pleasure, let me tell you, to work with Pat. We are on an issue here, for instance, where we have, at least he argues, a joint jurisdiction. [Laughter.] But, anyway, that is not up to me to decide.

Senator LEAHY. We certainly have a joint interest.
The CHAIRMAN. We certainly have a joint interest, which is the important thing. We have looked at his work, and it has been very helpful to us in our work. Pat, pleased to have you here.

Senator LEAHY. Thank you.

The CHAIRMAN. Do you have an opening statement?

Senator LEAHY. I would be happy to wait for the panel if you would like, Jim. Whatever you want. You are the chairman.
The CHAIRMAN. I think I will give you a chance perhaps after we question the panel.

Senator LEAHY. OK. Have you finished the panel?
The CHAIRMAN. We have finished. Why don’t you talk right now.

HON. PATRICK J. LEAHY, A UNITED STATES SENATOR FROM THE STATE OF VERMONT

Senator LEAHY. I will. Thank you.

The CHAIRMAN. I may have some questions.

Senator LEAHY. I do want to say how happy I am to be here. I mean, Jim and I have been partners on some of these things longer than most people have lived in Vermont, I guess, been married or something. [Laughter.] And so it is good to be here with good friends and fellow Vermonter. And when one of them is the chairman of the U.S. Senate Health, Labor, Pensions Committee, that helps even more.

Mr. Chairman, you and I introduced separate legislation, as you mentioned, last week to protect the privacy of medical records. We did this because we have a deadline. We have to have medical privacy legislation by August 21st of this year, and if we do not do it, then the Secretary of Health and Human Services does it. And I think this is far too important an issue to just go administratively. Jim, you and I support the same values, and I know in the end we are going to work together to pass a strong bill that is going to protect American families.

This is a time we have to worry about privacy. We see some states selling license photos. I must say I am very pleased that
Vermont declined to do that. Our leading chip and computer companies have built in a secret identifier so every time you use your computer they can find out where you went and nobody is told that that is in there.

In February, Jim, you and I held a hearing on home health care in Vermont. One person testifying brought the information of new regulations by Medicare and Medicaid called OASIS, and this requires home health agencies to gather very sensitive, personally identifiable information on all of their patients. The thing that struck me is they said on all of their patients, whether their claims are being paid by a government program or privately. I am very concerned about State and Federal governments compiling this kind of a personal database of sensitive personal health information, and I will be writing to Secretary Shalala to ask about this. It seems to be greatly overreaching to say we want this information on everybody whether we are paying for it or not.

It seems that both Big Brother and Big Business are trying to get at our most sensitive personal information, and I think that what it comes down to, if you have a medical record, you have a medical privacy problem.

Most of us think of our medical records being held in a manila file folder under the health care provider. When I walk in here I always stop by, there is a bill that is framed over here, it was a bill from when I was born at the Heaton Hospital. I hasten to add, Jim, the bill was paid [Laughter.] It is not a past due account, but I see Dr. McIntyre's picture on the wall, the man who delivered me. The fact is, it is not that way anymore.

When I was chairman of the Technology and Law Subcommittee of the Judiciary Committee I went into the privacy of medical records and new technologies. We found out good news and bad news. The good news is that we could do so much more for rural areas because of technology. The bad news is the information that is in there. One woman who testified told about going to a doctor for severe depression, and she was actually contemplating suicide. Got the proper medical care, obviously did not commit suicide, was helped with her depression. She was an immigrant. It is not the sort of thing she wanted to discuss in her family. To her shock her medical records were printed in the newspaper. Somebody took them. No laws were violated. And I asked her at the hearing, if you had known your records were going to become public, would you have sought medical help that you obviously needed? And she said, no, I would not have. And she might not have been alive. Instead, she is now a member of the House of Representatives and—U.S. House of Representatives and extremely well respected in there.

So we have to worry that if confidentiality is compromised it is going to deter people from seeking medical treatment or else stifle technological or scientific development. We should not fear the new technology. We should remember the technology is our servant and do not let it become our master.

In 1996 alone the health care industry spent an estimated 10 to $15 billion on information technology. Insurance companies want to know about your genetic makeup and health status so they can anticipate what diseases you will get later in life or whether they will
insure you to begin with. Pharmaceutical companies want to know what diseases you have so they can market their products to you. But the question comes down who controls that information, us or them? And if you are concerned about where it goes, that may reduce the willingness of patients to confide in physicians or other practitioners. Now I think the computerization if done right can ensure more privacy to individuals in the current system if we have strong medical privacy legislation, but if we do not, you have a great potential for embarrassment.

A few weeks ago The Wall Street Journal wrote about a company that is seeking the mother load in health data mining. The company wants to get medical data on millions of Americans to sell to any buyer, and currently there is no laws preventing that. That can be your information, any one of you, as part of that data gold mine.

Last year an article in the Washington Post described the story of a woman whose prescription purchases were tracked electronically by a pharmacy benefits management company two states away hired by her employer. Now when she used her card she got a discount. She thought that was great. What she did not know, all the sensitive information was being compiled, and her doctor was told that she would soon be enrolled by the company in a depression program and she would be watched for continued use of antidepression medication. She had been getting something for sleep and yet the company two states away who would interpret her medical records that she did not even know they had anyway, they decided that she was—better be watched out for because of her depression. This is wrong.

University of Illinois study found 35 percent of all Fortune 500 companies regularly review health information before making hiring decisions. And the health care providers are told they are constantly pressured to give out the most sensitive information to these companies or they will lose their own jobs. In genetic testing at the National Institutes of Health, 32 percent of the people who were offered in tests for breast cancer declined to take the free test. Why? Because they were afraid of who the information would be given to. Just stop and think about it. A test that could save lives, and they declined it because they are afraid of what might happen.

So, you know, the only—we can not let privacy slide the point. The only way for a person to ensure confidentiality is to avoid seeking medical care. We have got 159 days to correct this, and let me say I applaud you, Jim, for—you and your committee for being such strong leaders in this. I think we will put together a bill. I think we will have a bipartisan bill. I think we will have a strong bill. I think if we do not, health care and privacy will both suffer in this country. So thank you, Mr. Chairman.

[The prepared statement of Senator Leahy follows:]

PREPARED STATEMENT OF SENATOR PATRICK LEAHY

I am especially pleased to be here today with my good friends and fellow Vermonters, one of whom happens to be the Chairman of the U.S. Senate Health, Education, Labor, and Pensions Committee (HELP).

Mr. Chairman, you and I introduced separate legislation last week to protect the privacy of medical records. We support the same values and I believe, in the end, we will work together for passage of a strong bill to protect American families.
At a time when some states are selling driving license photos and information—which I am very pleased that Vermont has decided not to do—when our leading computer chip and software companies have built secret identifiers into their products to trace our every move in cyberspace without our consent, it is time for Congress to wake up to the privacy rights and expectations of all Americans before it is too late.

In February, Mr. Chairman, you and I held a hearing on home health care in Vermont. One person testifying brought up the issue of new regulations by Medicare, called OASIS, requiring home health agencies to gather very sensitive personally identifiable information on all of their patients.

I repeat—on all patients—whether their claims are being paid by a government program or privately. I am very concerned about State and Federal Governments compiling a database of sensitive personal health information. I have been in contact with the Administration about this issue and will soon be sending a letter to Secretary Shalala to get to the bottom of it.

It seems that both Big Brother and Big Business are vying to get at our most sensitive personal information.

The trouble is this: If you have a medical record, you have a medical privacy problem.

Increased computerization of medical records and other health information is fueling both the supply and demand for our personal information. We should not fear new technology, but unless we are vigilant, it will overwhelm our privacy rights before we even know what has happened. Technology is our servant; we should not let it become our master.

The debate boils down to a fundamental question: Who controls our personal information, and how freely can others gain access, buy and use it? We must not let privacy slide to the point that the only way for a person to ensure confidentiality is to avoid seeking medical treatment.

We have 159 days to enact a strong Federal medical privacy law. With the clock ticking toward the deadline set by Congress of August 21, 1999, let us act sooner rather than later. Mr. Chairman, I look forward to working with you on this very important issue.

The CHAIRMAN. Thank you. That is an excellent statement, Pat, and will be very helpful to us. We will be marking up our bill sometime in the not-too-distant future. I know we will be working with you as you have some excellent ideas in your bill. Ours—Senator Dodd and I put my bill in, so we have a bipartisan bill to start with. We are pleased to have you stay as long as you can.

Senator LEAHY. I would like to stay for a while if I might.

The CHAIRMAN. Tom.

Senator LEAHY. I know a number of the witnesses here. I look forward to hearing them.

The CHAIRMAN. On the issue of preemption, are you concerned that establishing a Federal ceiling would constrain State legislatures from enacting their own legislation?

Mr. KOCH. Yes. What would the point be if we have a federally uniform rule and State legislatures are preempted? It is a very complex subject matter, and that is why the bill has failed in the past two bienniums. It is the fifth year we are working on it, and what happens is that everybody has an interest in it and you end up with a real Christmas tree. And everybody has a special interest that, you know, they all come in and they want something added to it, and it gets very complex.

Federal uniformity I think is an easy answer, but I am not persuaded that it is the right one. I think states can often deal with their particular situations. But at the same time, we have a serious situation, and both you and Senator Leahy have outlined that very well, that needs attention at the Federal level. And I certainly hope that you do get a bill through here before August.
The CHAIRMAN. The legislation that I have introduced would give 18 months after enactment for states to establish stricter confidentiality rules. Would that be OK, or is it a problem for you?

Mr. KOCH. I do not think it will be a problem for Vermont because I think we are going to get this through if not by the time we adjourn this year, then by the time we adjourn next year. I mean, what happened here as far as I am concerned is I saw the bill fail in the last 2 years, I was not there the previous 2 years, and I said, what this bill needs is a shepherd, and so we are starting it in the House.

I do believe that the House Judiciary Committee is going to be working on it rather quickly. I think we will get it through the House this year, probably in the next few weeks, although the next few weeks on our agenda is extremely busy. Then it will go over to the Senate which is already working on it. Helen Riehle's committee, Senate Health and Welfare, has been looking at it, and so they are prepared for it. I am also told that it is the simplest version that has been around in the last 5 years because they did try to strip out some of the Christmas tree ornaments.

One of the things that we took out was the right to deny—in the health and welfare version we took out the right to deny mental health information to an individual because that individual may be injured in the doctor's opinion. And we heard from a lot of mental health advocates who said we have a right to see our own information. We do not need to be super protected from people who think that we cannot handle it. And our committee agreed with that, and we took that out of the bill. So it is a fairly stripped down version that we have, but we think it provides good protection for the people of Vermont.

The CHAIRMAN. Meg, does that provision in our bill, do you think, give you enough time to react?

Ms. O'DONNELL. I would agree with Representative Koch. Because Vermont has been looking at this issue so intensively for so many years, we are in a better position than some other states might be to react within that 18-month window, and that is, as I understand it, 18 months after the Congress would adopt this. So that is an even longer period. And I think for us that that would be ample time to accomplish something on the State level.

The CHAIRMAN. Mr. Palmer.

Mr. PALMER. I think that that does represent adequate time given all of the work that the legislature has done already, and I would reiterate my strong support for committing the states to do that because we in Vermont are very aware of the kind of confidentiality issues we have given the size of the State, and the number of people who live here and the closeness of all of us, that confidentiality around these issues is really important. And what is needed in Vermont may not be needed in California or Montana, but it is needed here. I think we are the best at being able to carve out the kinds of protections that we need locally.

The CHAIRMAN. Ms. Aranoff.

Ms. ARANOFF. I think I would defer to the legislators and regulators who would have to be meeting that 18-month deadline. If they feel comfortable, then I am sure that is accurate. I do know
that they have been working industriously for at least 6 years on this issue. So I am sure they could do that.

The CHAIRMAN. I guess this is more for Meg, but I think all of you can comment. What sanctions do you think are appropriate for those who violate patient confidentiality, and who should be responsible for enforcing those sanctions?

Ms. O'DONNELL. I am chuckling a little bit because I have seen all range of sanctions in the various bills before the Vermont legislature. The original version that was introduced in January of 1995 included I think the full range that I understand is in your bill which includes civil remedies, a civil action and criminal sanctions. And those were windowed down over time I think over a concern as to how much was needed to enforce.

I think the core of the sanctions has to be a civil right of action; that an individual who feels that he or she has been harmed by the use or misuse of their information have an opportunity to go into court and to fight it out there and get some remedy if, in fact, there has been misuse or abuse.

I think that it is also appropriate to look at maybe some kind of a higher remedy, whether it is a civil one enforced through the state's attorneys, or the Attorney General's Office, or perhaps even criminal sanctions, that are keyed in to more of the motivation behind misuse or abuse. Typically a lot of the model legislation we have looked at will include heightened sanctions in terms of heightened fines or even the penalty of imprisonment for multiple misuse, for misuse that is motivated by trying to get money or to harm somebody. And I do think that that is appropriate, and that is harder to put into a personal civil action.

So I would say that a broad range of sanctions is probably the most appropriate. I think we have to be cautious as to how you peg the crime or the civil remedy to what has happened because what you do not want to do is set up a system where a health care provider, for example an institution like Central Vermont Medical Center, is put under an obligation to do a broad range of things with their medical records to ensure confidentiality, finds itself in a position of having one employee who maybe mistakenly does something and then they find themselves on the hook for a large penalty or a large fine. I think you do have to be cautious that there is going to be probably some involuntary or some unknowing violations of this law, especially in the first couple of years as people are getting used to new ways of doing business.

The CHAIRMAN. Tom.

Mr. KOCH. I generally agree with what Meg had to say, and particularly her last comments about not coming down too hard on people who unintentionally make errors. We all make errors, and they are likely to continue. We do need requirements that organizations that maintain records develop procedures so people are aware of what is required. And I read in one of the other statements that I believe you will be hearing later this morning a suggestion that we have not had before our committee, but I think I will offer to the House Judiciary Committee when I appear before them on this bill and would suggest for your consideration, making it clear that a person who intentionally or maybe substantially violates someone's confidentiality in handling records, that that is legitimate
grounds for immediate dismissal. And I think if you make it clear that somebody's employment is at risk, you will get compliance. And it will also eliminate some litigation after a dismissal is imposed because, you know, if you do not say it, you will find a lawyer who will say it was wrong to fire that person for that reason.

The CHAIRMAN. Mr. Palmer.

Mr. PALMER. On that issue in particular, running a small organization that does have control over confidential information, we have confidentiality policies that protect those records. And if an individual staff person violates those policies, they are subject to immediate termination. I think that works in small settings, but when we are dealing with multistate corporations as Senator Leahy talked about and large employers, I think it is important to maintain the right of individuals whose confidentiality has been violated to civil action. And I think it is also important given the size of some of the employers and health care record organizations to also have criminal penalties involved because it would be easy for a client of Vermont Cares to take Vermont Cares to court because it is a much more even playing field. We do not have attorneys on staff waiting to handle any kind of complaint, but large corporations and large manufacturers and employers would have that kind of protection. And I think it is important to protect the individual's right. So criminal penalties I think are also important.

The CHAIRMAN. Ms. Aranoff.

Ms. ARANOFF. I would just like to add that I think it is very important that nominal damages, in other words, damages just for a violation, be allowed and attorney's fees be allowed for the simple fact that this particular area of harm is very difficult to prove. You know, even in some of the more egregious violations, the medical information that is sold to pharmaceutical companies for marketing, if you receive something in the mail saying, hey, there is a new treatment for X, Y and Z, which is something that you have, you receive that because your information was wrongly released, it is going to be hard to say how you were harmed by receiving notice of new treatment for your condition. Where is your damage? But I think that prosecuting civilly violations of these kinds of laws is a consumer protection function, and there has got to be some inducement for attorneys to take these cases when the damages are going to be either impossible to quantify or not very high. So I think attorney's fees are very important, and nominal damages would be very important.

The CHAIRMAN. Pat.

Senator LEAHY. Ms. Aranoff, if you got the brochure that says you can get 40 percent off on this generic, you have been buying this other, and we will argue that there is a benefit. Conversely, if it is promotion for a company and your employers look at this and say, well, they may be genetically disposed to breast cancer, prostate cancer, whatever else, so we are not going to promote it, or you are seeking insurance and similar—life insurance, similar things are given, though there would be no question in your mind what the damage was, I would assume?

Ms. ARANOFF. Yes. If you can establish you missed out on a promotion when you were hired initially, you would have a good case for damages.
Senator LEAHY. And in sensitive health information, and it could be mental health; it could be reproductive health information; Mr. Palmer, it could be information you handle at Cares, the bill that I have put in specifically allows patients to restrict what kind of information can be released from their records. Is that something that you think makes sense?

Mr. PALMER. I think it is absolutely essential for individuals to have control over what pieces and parts of their medical records are made available. I think the system has to be structured in a way that by saying that I do not want my HIV status disclosed, by saying that I do not want it disclosed, that I am not disclosing it in that form itself. So I think that the—that how the program is structured, how your control over your medical records is structured has to be in a way that protects your confidentiality right from the very beginning.

Senator LEAHY. It is not directly related, but I think of the number of times in our military, for example, when somebody did not seek help for alcoholism, partially because they are afraid they would not get their promotion. I saw cases similar to this in law enforcement when I worked in law enforcement, and it bothers me. Tom, I was glad to hear your comment about needing a guardian angel or shepherd on this, and I think it is absolutely important. I mean, everybody—if you ask 12 people walking out of here, do you think you ought to have some privacy in your medical records, of course they will all say yes. It is a long way from there to a bill that is workable, enforceable and makes sense. And one of the debates we will have in the Congress will be on the question of preemption and how much a Federal law might preempt State law. It is something that when you come from a small State like ours you are, I think, particularly sensitive to.

How do you feel about a law which—a Federal law that sets a basic floor but allows a State at any time, either before or after, to set a tougher law on disclosure if they feel it is necessary to—if the legislature of that State feels it is necessary to protect the people of that State?

Mr. KOCH. I was the first member of the panel to speak, and I spoke before you got here, and essentially that is the concept that I did endorse, that we have a Federal floor and that states be allowed to extend greater protections. In answer to Senator Jeffords' specific question, I responded essentially that I think 18 months is probably sufficient time for us to act on this bill. But things change over time, and the situation may come up that Vermont wants to extend some protections to 5 years from now, and if the Congress does not act, then Vermont would be powerless to act. So my statement was directed essentially to the issue of Federal preemption, and I would prefer that we set a minimum Federal standard and allow the states to extend greater protections to their citizens if—

Senator LEAHY. You will not be surprised if I tell you I agree with you completely on that one, and that is written out well in the bill. I wish I had heard the earlier part of it. I was shoveling out of Middlesex a little bit earlier this morning, although this—when I watched the news this morning, Jim, they are saying light snow, accumulations of no more than two to four inches. In Washington
when they are expecting two to four inches they break into the news every 2 minutes with trailers. [Laughter.] They have got people out, they film the first snowflake coming down to get the appropriate level of panic on the people. [Laughter.]

Thank you very much.

The CHAIRMAN. On the other hand, Mr. Palmer, just to get to the reason that we need to have transfer of information a little bit into the equation here. For instance, suppose you have HIV and you do not want that disclosed, but non-disclosure could mean getting some pharmaceutical treatments, drug treatments, which might be—create an adverse drug reaction. How do we handle that problem?

Mr. PALMER. Well, I think that one of the realities we have learned through the course of the epidemic is that individuals living with life-threatening diseases have themselves a responsibility to take care of their health care. And if we were to just rely on the system, we are not going to protect ourselves. So I would err in favor of letting the individual have responsibility for keeping themselves informed, not using information about them to inform them.

I think that the ownership of the information needs to be the patient, not a system. If I have a condition that I am aware of, I am responsible for tracking the treatment that should be available to me. I do not want big pharmaceutical companies, I do not want big medical providers, I do not want my employer to provide me information about the best course of treatment for me.

The CHAIRMAN. That is a partial answer. What I am thinking of is this: suppose you are taking certain medications for HIV and that is not disclosed, at the same time, another physician is treating you for something else without knowing about the HIV drug. A certain drug combination could cause an adverse, maybe fatal reaction; how do you protect against that?

Mr. PALMER. Well, I think that there are a number of safeguards. Again, in working with your primary care physician, it is your responsibility along with the physician to—to identify your entire course of treatment and what medications you are in fact taking. I do not—again, I would err on the side of the individual patient and physician to have that responsibility to track what interactions might or might not occur, not to have some electronic system controlled by somebody else knowing more about me than I know about myself advising me what kind of course of action I should be taking.

The CHAIRMAN. Ms. Aranoff, what single issue involving confidentiality is of most concern to the mental health community? There are many of them.

Ms. ARANOFF. There are so many of them. The issue that comes to our attention most in the office is the issue of disclosures to family members and people that the person is associated with without any kind of release and often against the person’s will. In fact, it is the policy of many of the community mental health centers and it is the policy of the State Department of Mental Health to include family members on people’s treatment teams, what they call treatment teams, and to—and they have a memo out that says—and this does not violate confidentiality laws.
And I worked with a client who was being discharged from the State Hospital into the community, and the community based provider would not take her—would not provide the services to her, services that the state's obligated to provide, unless she would allow her family members in on the conversation and in on her treatment and to have access to her treatment. And that kind of coercion to violate confidentiality and that kind of routine disclosure to family members, without the bill that is before Senator Koch's (sic) committee, Vermont statutes specifically allow health care information about people who have been committed to be released to family members. And very often it is the patient's position that it is the family members that are either responsible for their conditions or responsible for them having been in the State Hospital to begin with, and they are the exact people that they do not want in on their information. So I would say that that is the issue that we receive the most complaints about.

The CHAIRMAN. Thank you. I just want to make a comment on the other side of the question which we have to be aware of, and that is the importance of the ability to trade information in outcomes research to determine how we can better treat patients. We have to weigh that balance, and it is not easy to do.

But also as Senator Leahy, I think, pointed out earlier, with modern technology, information transfer can be done very selectively. You can get the core information you need in order to improve treatments and to determine how well treatments work, in very sophisticated ways, by passing data that are not relevant.

One of the things we are considering at the Federal level in the large amount of information transfer occurring now. This fall, we are probably going to be linking up with Europe concerning all sorts of treatments being used around the world now, and the outcomes of various treatments moreover to encumber the free flow of useful information between hospitals across State lines could actually have adverse effects on the health of our Nation's citizens. So it is not an easy issue, and you all know that.

I assume, Tom, you are aware of that. Let me ask you this about the transfer between New Hampshire and Vermont for instance. How are you handling that?

Mr. KOCH. I believe that—well, that is one of the problems that I would hope that the—you would take reins on. It is—once the information goes across the State lines, we do not have true jurisdiction, and it is difficult. We have enforcement problems, although we probably have a minimum context jurisdiction in a civil action. And I do not know what New Hampshire is doing, frankly. I really cannot give you a good answer.

The CHAIRMAN. OK. I would draw from that that there is a role for the Federal Government to play—

Mr. KOCH. Absolutely. Absolutely.

The CHAIRMAN. —in some sort of uniformity and in some system to maximize the beneficial utility of information.

Mr. KOCH. Or as I suggested, giving effect to the originating State's law which might have a definitional problem.

The CHAIRMAN. Senator Leahy.

Senator LEAHY. No, I have nothing further. Thank you.

The CHAIRMAN. You are the next panel.
Senator Leahy. And I have already spoken. [Laughter.]

The Chairman. All right. Thank you very much. Very helpful. I would like to reserve the right to ask you further questions in writing if you do not mind, because this is an extremely important issue for all of us. And my number-one concern is Vermont. So I do not want to do anything at the Federal level which will inappropriately interfere with your work, and I want especially to keep in close touch with you, Tom, as you progress. That's also true of the other members of the panel. Thank you very much. Very valuable testimony.

Our next panel is Leigh J. Tofferi, director of government and public relations at Blue Cross & Blue Shield of Vermont. Leigh, if you will come forward. He currently develops public policy and also has primary responsibility for liaison with State and Federal legislative bodies, governmental agencies, news organizations, and the public. Wow.

Prior to joining Blue Cross/Blue Shield, Mr. Tofferi served four terms in the Vermont House of Representatives. Thank you for being here.

I would also like to introduce Ms. Anne Cramer who is a partner in the Burlington law firm of Miller, Eggleston & Cramer, Ltd. In her private practice she represents hospitals, nursing homes, physician groups, and various private groups in Vermont. Today she is representing the Vermont Association of Hospitals and Health Systems. Among her professional accomplishments, Ms. Cramer was one of the four original drafters of proposed comprehensive legislation addressing the collection, use, and disclosure of confidential health care information introduced in the Vermont legislature in 1995.

A welcome to you, and we are fortunate to have you here today. Appreciate it.

I am next pleased to introduce Madeleine Mongan, who is here today representing the physicians, residents, and student members of the Vermont Medical Society. Welcome, to you.

Please proceed.

STATEMENTS OF LEIGH TOFFERI, DIRECTOR OF GOVERNMENT AND PUBLIC RELATIONS, BLUE CROSS AND BLUE SHIELD OF VERMONT, MONTPELIER, VT; ANNE CRAMER, VERMONT ASSOCIATION OF HOSPITALS AND HEALTH SYSTEMS, BURLINGTON, VT; AND MADELEINE MONGAN, COUNSEL AND DIRECTOR OF GOVERNMENT RELATIONS, VERMONT MEDICAL SOCIETY, MONTPELIER, VT

Mr. Tofferi. Thank you, Mr. Chairman. I am testifying today as a representative of the state's largest private health insurer—

The Chairman. Excuse me. I want to ask you all to speak up because we have an overflowing crowd here which is kind of—so aim at them and speak to me. [Laughter.]

Mr. Tofferi. I am representing the state's largest private health insurer and the only health insurer domiciled here in the State of Vermont. Blue Cross & Blue Shield of Vermont provides health care benefits to approximately 190,000 Vermonters in every category of health insurance: The individual market, small and large group, Medicare supplemental benefits, the state's safety net pro-
gram and for Medicaid recipients through a contract with the State. I thank you, Mr. Chairman, for the opportunity to express our views about this important issue.

Blue Cross & Blue Shield of Vermont considers the confidentiality of the subscribers' medical records to be of prime importance. The plan has a strict policy in place to protect subscriber privacy but which also allows an efficient process to administer health benefits and pay medical claims.

Currently our plan certificates of coverage include a provision clearly stating the plan's right to obtain the information necessary to administer benefits and resolve grievances. By accepting this contract, the subscriber agrees to allow the plan to obtain the necessary information to process claims and perform functions such as utilization review. In return, the plan agrees to pay for claims under the terms of the contract.

This agreement provides the plan with sufficient authority to gather the information necessary to perform the functions we all expect of a health care plan. The plan recognizes its responsibility to protect this information and employs tight controls on information it obtains.

The release provision in the certificate of coverage includes our subcontractors as entities that may receive medical information. However, the plan protects the use of that information through provisions in its own contracts with those subcontractors requiring them to comply with its protections for medical information.

We also have a strict employee confidentiality policy in effect. This policy covers internal communications of health care information, external communication in response to customer service inquiries and external communications of statistical health care information. This policy also covers retention and disposal of records and any other instances governing the use of medical information. All employees are briefed on the confidentiality standards of the plan and sign a statement acknowledging that they understand the plan's confidentiality policy.

We will only release medical information to the patient or person specifically authorized by the patient and then only with a release from the attending physician or medical provider. The plan may provide a spouse or family member with benefit information but will not release medical information without written authorization.

Although our current standards have worked well, we support the concept of a clear and concise approach to medical confidentiality. We have worked with the Vermont legislature as it has considered these issues over the past few years, and we also look forward to working with you.

In that regard, we have adopted three principles to guide us when discussing this topic. First, legislation in this area should balance the protection of consumers' personally identifiable health information with the needs of the health care industry to deliver the best possible service. Second, each person or entity in the health care chain must be required to fully participate and assist in ensuring the protection of individually identifiable information. Third, legislation for access, use and disclosure of this information must be clearly defined to avoid misapplication, misuse of the law.
We have also identified some key issues that could adversely impact our ability to function if not appropriately addressed. Authorizations for disclosure and use of protected health information for health plan functions should not be required. For health plans this might include payment, credentialing, fraud detection, disease management, utilization review, etc. If, however, authorizations for health plan functions are required, a single authorization at the time of enrollment for these purposes for the life of the relationship between the insured and the insurer should be the rule. Preferably, the law would provide by statutory authorization or deemed authorization to be granted at the time of enrollment.

Special protections should not be created for sensitive categories of health information, for example disease-specific information. Any legislation should protect all types of individually identifiable health information equally including genetic information.

Individuals should be allowed to review and copy their protected health information from entities that collect, transmit or process this information, but health plans should be allowed to require a provider release before issuing records through an insured member.

Any new confidentiality law should recognize and appreciate that health care is local. Any enforcement structure established by any Federal confidentiality law should, wherever possible, link national and State, local resources.

Finally, any penalties should clearly define the behavior that constitutes a violation. Further, they should be consistent with the threshold/intent standards and penalties established in current law. The penalty structure should allow for the opportunity to take corrective action and should establish intermediate sanctions. Enforcement should not include private right of action or punitive debarment from Federal programs.

We look forward to working with you on this important issue, and thank you for giving me the opportunity to express our views.

The CHAIRMAN. Thank you very much.

Ms. Cramer.

Ms. CRAMER. Thank you, Mr. Chairman. I am here today and testifying on behalf of the Vermont Association of Hospitals and Health Systems. We have 16 member hospitals in the State of Vermont, 14 are acute care institutions; in addition, the Brattleboro Retreat and the Veterans Administration Center.

This is an issue that has been very important to us. We have been long looking to have a void filled. As a slight digression, when I moved to this State in 1985, my law partner, Marty Miller, said, ah, some people would like us to give a seminar on medical records. Why don't you do it. It will be a good way for you to get to know the State. So I started doing some research and found out that I was really going to be making a lot up, that the medical records law on the books did not cover nearly any of the scenarios that had been painted to me as being important questions to the clients that we were working with both in hospitals and patient offices.

We thank you on behalf of the association for this opportunity to testify and work with both of you. We are grateful that this is a priority, and we have—also want to just make sure that there is not any misunderstandings I think as to our efforts at the State level as well as our really strong desire that there be uniform pre
emptive Federal legislation that defines the scope of patient rights as well as the obligations of all of us who either generate, store, transmit or use confidential health information.

Since 1993 there has been a pretty extensive effort at the State level to draft legislation. The Hospital Association, as it was known then, was very much in the middle of it trying to make sure that we fill this void, that we define what information is actually covered; that there is standards for what is an appropriate authorization; we define circumstances where it is appropriate to have information disclosed without an authorization; that we establish firmly the rights of the individual to copy their records, to access their records, to correct or seek to comment on them.

Additionally, as we have moved into the electronic age, the security standards are something that we are all looking to have uniform so that we make sure we are designing these expensive information systems in the right manner. And, indeed, we also believe there should be serious penalties for wrongful disclosures.

We have worked at the State level and we still continue to work with Representative Koch, and we hope that, indeed, a bill can be passed at that level. This is a difficult issue, and as much as we are hopeful that between now and August you can accomplish your work, we feel that this is a race to at least get the void filled. We would, however, prefer to have Federal legislation, and we would like it to be preemptive, both as a floor and a ceiling.

The hospitals and doctors today work in an interstate environment. They seek tertiary care out of State. Lab results, lab testing goes out of State. We are back and forth across State lines constantly for the care management of patients. As much as we would like to keep this as an intimate Vermont relationship, it is not. And our biggest concern is that if you put the type of tough penalties in place that should be in a law of this caliber, that if we have states with varying standards, all for good reason and all, I think, trying to do the best things for their own population, that we are going to be in a situation where we cannot comply, that we will be violating the law. We will be in a catch 22 where we violate a New Hampshire law but it complied with perhaps Federal and Vermont.

So from this standpoint I think this is an area that I know there is a lot of—there is a lot of hard work that you do not want to deny at the State level. On the other hand, health records mean a great deal in a lot of different aspects of society and our ability to continue to deliver and improve the quality of health care that we deliver, that this really is something that needs Federal preemption.

The other area that I would like to just address briefly today is we do not believe there should be special rules for special diseases. We are very sympathetic with the AIDS community, but as we have gone on and talked with our patients, there needs to be one standard. And there is no question that all of us need to improve our culture of confidentiality, so whether or not it is a pregnancy or an HIV, there is a very strong standard that this is personal information and it needs to be protected and respected.

We also are weary and really do not support any type of effort of a patient block from one provider to another. Although for some areas you may not think that the mental health notes may be relevant to a primary care physician, there is a lot of different steps
in health care delivery, a lot of different scenarios we cannot always anticipate. And to have some artificial blocks put in place both makes it very difficult to manage the record and it also creates potential where we are not going to be able to give the care that is appropriate.

So I thank you, and I look forward to working with you.

[The prepared statement of Ms. Cramer follows:]

**PREPARED STATEMENT OF ANNE CRAMER**

Mr. Chairman, I am Anne Cramer, a lawyer with the law firm of Miller, Eggleston & Cramer which serves as legal counsel to the Vermont Association of Hospitals and Health Systems ("VAHHS"). VAHHS has sixteen member hospitals which include fourteen acute care facilities, the Brattleboro Retreat and the Veterans Administration Center in White River Junction.

On behalf of VAHHS, I thank you for this opportunity to testify on the extreme importance of enacting strong, uniform and preemptive federal legislation defining the scope of patient rights and the obligations of all who generate, store, transmit or use confidential health information. Health care delivery at our member hospitals necessitates the involvement of interstate providers and payers and requires that health information be regularly transmitted across state lines for such purposes as obtaining tertiary care, lab testing, care management, quality assurance review and long term care services. Federal confidentiality legislation must set a national standard, which is both "the floor and the ceiling", for protecting patient privacy while ensuring that information is available for patient treatment.

Currently, neither federal nor Vermont law addresses the confidentiality of health care information in an appropriately comprehensive manner. In particular, Vermont's present law, 12 V.S.A. Sec. 1612, consists of a simplistic statement of the patient's privilege against unconsented disclosures of health information which is virtually impossible to apply to the hundreds of scenarios where access to health information is sought for a multitude of purposes beyond the delivery and financing of health care services.

To address longstanding concerns relating to the privacy and management of health information, in late 1993, VAHHS participated in an effort initiated by an organization then known as the Vermont Health Information Consortium ("VHIC")—a private nonprofit organization funded by a grant from the John A. Hartford Foundation—to explore the development of a statewide health information system. As part of that process, VHIC conceived a set of fundamental health information privacy principles to serve as a basis for the development of legislation for introduction and adoption by the Vermont General Assembly. I was one of four individuals who worked to draft comprehensive health information legislation for Vermont on the basis of these fundamental principles. The first bill was introduced in the Vermont legislature in January of 1995 and sought to accomplish the following objectives:

1. Define the scope of health related information entitled to confidentiality;
2. Provide a standard for an appropriate written patient authorization for disclosures;
3. Establish the circumstances under which health information could be disclosed without a patient's authorization;
4. Establish the rights of an individual to obtain copies of his or her own medical records or seek to correct or comment upon them;
5. Establish standards for how health information should be maintained, secured and retained by health care facilities and providers;
6. Establish damages and/or penalties for violations of this legislation.

This proposed Health Information legislation has received extensive review, revision and debate in every year of each of three biennial sessions of the Vermont legislature since 1995. Today two comprehensive bills are under current consideration.

VAHHS and its member hospitals have been very supportive of state legislation to fill the existing statutory void and patchwork of court precedent. As health care professionals and institutions are required to better manage the cost of delivering health care, electronic information systems have increasingly been utilized to collect, sort and transmit pertinent health information. While these information systems offer greater opportunities for delivering a higher quality of health care, they create new problems for potential misuse and breaches of security. Clear standards for the operation and security of such systems is imperative.
It is our preference that federal, rather than state, legislation be enacted so that all patients are protected uniformly across state lines since Vermont's health care delivery system delivers care to residents of other states and is extensively integrated with providers and third party payers in other states. Federal legislation must be designed with the following tenets:

1. It must preempt state law so providers can rely on uniform standards and are not faced with trying to comply with potentially inconsistent or conflicting requirements. In this regard, federal confidentiality legislation must be the "floor and ceiling" of standards with no more than an occasional specific exception, perhaps for public health reporting.

2. The requirements must consider the realities of our multifaceted health care delivery systems and not serve to prevent the appropriate flow of health information between individuals treating a patient. The notion of a patient "block" preventing certain providers from accessing and using health information during treatment is untenable. Health information must be fully available for diagnosis and treatment services. Special disclosure rules for special diseases also has the potential of prohibiting the appropriate flow of information to health care providers when it might be most needed. It is imperative that there be a single standard of confidentiality.

We thank you for this opportunity to set forth our views and look forward to participating in a more detailed discussion with the Committee focused on the goal of protecting patient confidentiality and promotes the efficient delivery of high quality health care through a uniform national standard.

The CHAIRMAN. We look forward to working with you.

Ms. MONGAN. Thank you. Good morning. Senator Jeffords, Senator Leahy, my name is Madeleine Mongan. I am counsel and director of government relations for the Vermont Medical Society, and I thank you for the opportunity to speak to you on behalf of the approximately 1400 physicians, medical students and—physicians residents and medical students who are members of the medical society. And thank you for coming to Vermont and holding this hearing and for making this a priority issue for both of you both introducing legislation.

My remarks are going to be based on a formal position that was adopted by the Vermont Medical Society Council in October of 1995. We have been working on this issue for a long time, and we are excited to see it, we hope, making progress on both the State and the Federal level.

Because the physician-patient relationship is based on trust, the medical society believes that the confidentiality of communications within this relationship is essential to high quality medical care. Privacy of medical records is ingredient to this ability to trust your physician or your other health care provider. And in order for physicians to provide the best and most appropriate medical care, patients must feel that they can disclose to their physicians personal facts and information that they would not want others to know. Without assurance of confidentiality, patients may not provide the information that their doctor needs to properly diagnose and treat the patient.

Now the second factor leading to the importance of this issue at this time is, as you know, the evolution of electronic medical data. This has intensified concerns, although I agree with other speakers that have said this may create the ability to provide greater confidentiality rights. It has intensified concerns, particularly with the linking of various databases.

The Medical Society believes that Vermont's patients will benefit from the adoption of Federal legislation and State legislation that protects the confidentiality of medical records. We have some cri-
teria for this legislation. It should clarify patient and provider un-
derstanding of their rights and responsibilities. As Anne said, when
we first began looking at this issue, it is—in Vermont the laws are
a mish-mash or a hodgepodge.

We are looking—also consistent with Ms. Cramer’s statement, we
are looking for a law that will create consistent nationwide stand-
ards for confidentiality of medical records that will apply in all
health care settings. We are hoping that confidentiality will be able
to be protected with minimal increases in bureaucracy for physi-
cians who are burdened at this time by paperwork and bureau-
cracy, and we are also hoping that confidentiality legislation will be
sufficiently flexible to adjust to future changes, both in medical
care delivery and in record keeping processes and electronic data.
The legislation should include a fair time frame for implementation
and should be premised on the assumption that a law governing in-
formation practices cannot guarantee confidentiality. Protection
must—real protection has to result from creating, as others have
said, a culture of confidentiality and educating record keepers, pro-
viders and others who deal with this information on the—both the
clinical and the ethical importance of privacy.

Now I will speak briefly on the preemption issue. We, as I said,
are looking for a consistent national confidentiality standards, and
we think health care is not confined within a set of State bound-
aries. Vermonters travel to our border states regularly. They travel
to other states to get specialized medical care. And we have physi-
cians who are practicing—southeastern Vermont is one example
where I know of some physicians who have offices in three states,
in Massachusetts, in New York and in Vermont. And they may be
seeing patients from any of these states in any of these offices on
any 1 day based on the schedules of the patients and the physi-
cians.

We think that if there are different confidentiality requirements
imposed by different states, potentially they could be inconsistent
and this could create a tremendous burden for physicians in other
health care facilities. Medical records need to move with patients
to ensure quality health care and should be subject to the same
and I would say a high level of protection regardless of whether the
patient is in Montpelier getting treated for a sore throat or is in
Pittsburgh receiving a heart transplant.

Others have mentioned telemedicine which is another cross State
lines issue which is important for rural health care in Vermont.

And final point on this is the various number of Federal and
State programs that physicians participate in with potentially in-
consistent standards. To the extent that the standards can be the
same for all programs, that would be I think helpful to both pa-
tients and to physicians.

Now I would like to digress from my written statement for a
minute to offer a few comments on S. 578 which I received Satur-
day.

The CHAIRMAN. Yes. Please do.

Ms. MONGAN. OK. We are very supportive in general of the legis-
lation. We are very happy to see it introduced. In Title 1, Subtitle
A, and this goes along with my statement, we are very supportive
of the provisions governing patients access to their records. We also
support an exception to that access in situations where there is
danger to health, life, and this is from the bill, the language in the
bill, the exception in the bill, if there is danger to health, life, phy-
sician's safety or if substantial mental harm would result. I think
it is important to have that exception. We have a similar exception
in Vermont law and other states which allows physicians to inform
victims—identified victims if there is a risk of harm—a threat of
a risk of harm to that identified victim. So I think this is a similar
kind of exception that we need that will be used very rarely but
needs to be there as a safety valve. And I know that is important
to physicians in Vermont.

We also support the ability of patients to amend their records.
We have a concern about the 30, 45 and 60-day time lines, and
that concern is that there should be a good cause exception for
those rare cases when these time lines could not be met.

And as far as the notice of confidentiality practices, this imposes
some bureaucracy for physicians, but it does say that the Federal
government is going to create a model notice which would be help-
ful and that physicians can either choose to put it on the wall or
to hand it out or provide it to patients which gives them an option.
So we are generally supportive of that.

In Subtitle B we support the safeguards but have a very small
concern—the safeguards require the person keeping the record to
maintain the accuracy of the record, and that might not be possible
in all cases. There could be some inaccurate information that gets
into the record from—maybe the patients do not provide the history
quite correctly. And so we are concerned about that becoming a
mandate in view of the penalties.

On the disclosure tracking provision, we understand that the de-
tails of this will be promulgated by rule, but our concern is that
the tracking be very simple and straightforward, particularly for
disclosures, okay, among physicians. I will be able to submit the
rest of these comments to you.

[The prepared statement of Ms. Mongan follows:]

PREPARED STATEMENT OF MADELEINE MONGAN, ESQ.

Senator Jeffords, Senator Leahy and members of the committee, my name is Made-
eleine Mongan, I am Counsel and Director of Government Relations for the Vermont
Medical Society. On behalf of the approximately 1,400 physician, resident and medi-
cal student members of the Vermont Medical Society, I appreciate the chance to
comment on the issue of medical records confidentiality and thank the committee
for holding this hearing in Vermont and devoting the time to discuss this important
issue. My remarks are based on the Vermont Medical Society's Position on Access
to Confidential Health Care Information which was formally adopted by the VMS

Because the physician-patient relationship is based on trust, the Vermont Medical
Society believes that the confidentiality of communications within this relationship
is essential to high-quality medical care. In order for physicians to provide the best
and most appropriate medical care, patients must feel that they can disclose to their
physicians personal facts and information that they would not want others to know.
Without assurance of confidentiality, patients may not provide the information nec-
essary for proper diagnosis and treatment to their physicians.

The evolution of electronic medical data has intensified concerns about access to
patients' confidential medical information. Concerns regarding the transmission and
aggregation of electronic data are amplified by the practice of linking information
across data bases.

The VMS believes that Vermont's patients will benefit from the adoption of fed-
eral legislation that protects the confidentiality of medical records if the legislation:
• clarifies patient and provider understanding of their rights and responsibilities;
• creates consistent nationwide standards for confidentiality of medical records that apply in all health care settings;
• enhances confidentiality with minimal bureaucracy;
• is sufficiently flexible to adjust to future changes in the medical care delivery system and in consent and record keeping processes;
• establishes a fair time-frame for implementation of any significant changes in current access laws; and
• is premised on the assumption that a law governing information practices cannot guarantee confidentiality: such protection will result from educating record keepers on the clinical and ethical importance of privacy.

Consistent National Confidentiality Standards

Health care is not confined within any one set of state boundaries. Vermonters travel regularly to health care professionals practicing in our border states—New Hampshire, New York and Massachusetts—to receive health care. They travel even farther to receive specialized services. Likewise, many physicians practicing in Vermont treat patients in other states; some have offices in two or, in the case of southeastern Vermont, even three states. In view of the mobility of health care delivery, the imposition of different confidentiality requirements, potentially inconsistent, created by different states, on physicians and health care facilities could create a legal and bureaucratic nightmare. Medical records need to move with patients to ensure quality health care, and should be subject to the same level of protection regardless of whether the patient is being treated in Montpelier, Vermont for a sore throat or receiving a heart transplant in Pittsburgh, Pennsylvania.

An increasing number of physicians in Vermont are participating in telemedicine initiatives, that make it possible for them to consult and treat patients in remote and often rural locations, both inside and outside of Vermont. Additionally, regardless of where they provide care, Vermont physicians participate in a number of federally operated or funded programs that each have their own rules and standards. Because of the increasing complexity of the health care system, a clear set of consistent confidentiality rules that apply across state lines, across all federal and state programs and across facilities is of critical importance to physicians, and the future of the health care system. Confidentiality rules should not impede the efficient delivery of health care.

Purpose of the Medical Record

The primary purpose of the medical record is to provide a reliable tool for clinical diagnosis and treatment of patients. Medical records must be accurate and complete in order for patients to receive high quality medical care. While patient privacy is of vital importance to high quality medical care, this need for confidentiality must be balanced with the need to ensure that the medical record is accurate and complete. In that regard, the VMS cannot support including a provision in the law that would allow patients to limit or block the transfer, among health care professionals, of information that is necessary to provide medical treatment. Such blocks on the transfer of records could lead to duplication of tests or treatment. More important, because patients are not necessarily aware of the consequences of limiting access, limiting information could result in harm to patients through adverse drug interactions or unforeseen complications.

Provider to Provider Record Transfers

It is a physician's duty ethically and legally to provide medical information on a patient to another provider when the information is reasonably necessary to provide health care services to that patient. A physician's failure to provide a record to a subsequent physician or hospital constitutes unprofessional conduct which could result in loss of license. While such transfers do not require the explicit consent of a patient, implied consent is presumed to exist when a patient seeks the services of another physician or hospital. When consent is not clearly implied physicians customarily seek written permission from the patient.

The VMS can support a patient's right to ask their provider to refuse to disclose certain parts of their record to another provider only if:
• such a block on the record can be structured to insure that the receiving physician is aware of the block and can freely elect not to treat the patient;
• the licensing board acknowledges that such a block is an exception to the implied consent regarding transfers to subsequent physicians; and,
• the physician is protected from medical malpractice suits arising out of a block on the record.

Sensitive Information

Patients must be treated as whole individuals. Sensitive information about all aspects of the patient's history and treatment that is necessary for treatment must be available to physicians. Legislation should ensure that the confidentiality of all
health information is protected. Creating a different level of protection for certain health care conditions implies that all information is not subject to equal protection. Information concerning sensitive conditions such as HIV or mental health can be necessary for treatment purposes and should be maintained as part of a patient's medical record. Segregating sensitive information that is necessary for treatment could impede quality treatment. Likewise, a separate release form should not be required for certain types of information. A provision that limits release to information necessary to accomplish the purpose of the disclosure is sufficient to ensure that extraneous information is not released.

Paperwork Burden

Physicians are already overburdened by paperwork. Any confidentiality bill should be designed to reduce rather than increase bureaucracy. While the VMS supports the patient's right to know when a record transfer has occurred, it does not support creating a bureaucratic tracking system, particularly for small rural providers. Physicians should be able to document disclosures, particularly to other providers, in the patient's chart, and should not be required to maintain separate and detailed documentation of disclosure. This is an area in which legislation must be flexible enough to await the time when the technology is in place to track records movement without undue red tape.

Patient Access

Patients should have access to a copy of their medical records to assist them to understand their health status. Any limitations on the patient's ability to obtain the actual record should be narrow and in the patient's best interest. If it is determined that access to the record will endanger the patient or others, the patient must have recourse such as:

- receiving a summary of the record;
- obtaining a second physician's opinion;
- appealing the decision.

The statute must be clear that the patient may obtain all of the record regardless of whether the content was:

- authored by their own physician or other staff;
- from the hospital;
- from a previous physician; or
- inclusive of comments made by a third party—for example a family member.

This change in access practices must be carefully implemented with an effective date which gives providers sufficient time to conform to new procedures. The law should be clear that records collected prior to the effective date are grandfathered allowing providers the option of summarizing records prepared previous to the effective date.

Access to a patient's record includes access by a parent or guardian to a minor's record; however, minors must have the right to block parental access to their records if the records contain information on treatment or procedures to which minors have the legal right to consent.

Record Ownership

While a medical record is about a patient and the patient must have open access to that information, the owner of the record must be the person who is primarily responsible for:

- ensuring that the record is accurate and current;
- deciding what recording methods are used written, taped, video recording, or computer retrievable;
- implementing security measures;
- establishing consent and disclosure protocols;
- preserving the record;
- ensuring that the record is maintained in a manner which enhances appropriate medical care.

The VMS advocates for legislation which is centered around the patient-physician relationship.

Amending The Record

After reviewing medical information, a patient may disagree with the physician's conclusions or may have a factual correction to make. In either case, the patient should have the right to add a notation. The record owner may also disagree with the conclusions of a consulting physician or may find that a statement made as fact was later found incorrect. Neither the record owner or the patient may erase or cross out the original record, but both may make notations. Notations should be reasonably brief.

Access to the Record Without Authorization
There are a number of instances under existing law by which a medical record should be made available to someone other than the patient or the owner without consent assuming the disclosure follows reasonable confidentiality protocols:

• in a peer review process;
• in a medical emergency to a provider or family member if the patient is incapacitated;
• to meet infectious disease or cancer registry reporting requirements;
• when a state agency is fulfilling its obligations to investigate insurance or workers' compensation claims.

Access by Researchers
At a minimum, legislative change is needed to assure that researcher's access to confidential information used for research purposes is controlled by an Institutional Review Board which shall consider whether the confidential records can be traced back to the patient; the degree to which the research process or outcome could have an adverse affect on the participant; and federal laws limiting access to mental health and substance abuse records.

Release Under Court Order
The law should make clear that a physician shall not transfer a medical record after receipt of a subpoena until either:

• the patient signs a consent form; or
• a court order is issued.

In that event, a physician shall release relevant records to the appropriate parties in a timely fashion. However, a physician must retain the right to appeal a court order on the basis that release of a complete copy of the medical record would be injurious to the patient.

Consent Forms
The law should establish minimum requirements for the content of consent forms including the following:

• the need for and proposed use of such information;
• a statement indicating specifically the type and extent of information to be released;
• a statement that such information will not be given, sold, transferred; or in any way relayed to any other person or entity not specified in the consent form or notice without first obtaining the individuals additional written consent on a form stating the need for the proposed new use of such information or of the need for its transfer to another person or entity;
• the duration of the consent; and
• the patient's-signature attesting to understanding the implications of the consent.

Security
The record owner is responsible for keeping the records secure from inappropriate access and must follow appropriate protocols including stringent security measures applicable to all staff whether the records are kept on paper, tape, video or computer. Patients should have the right to know what form of record keeping is employed by their provider and the security measures followed to maintain privacy.

Where confidential health care information is in computer-retrievable form such information shall be subject to the following minimum measures:

• security clearance available only to authorized individuals;
• identified individual responsible to maintain security procedures; and
• prohibition on contractual limitations of liability for breaches of confidentiality.

Penalties
Civil and criminal penalties should be sufficiently large to deter intentional and knowing violations of this Act. Stiff penalties should also apply to anyone who obtains an individuals confidential health care information through the commission of a crime. However, we are concerned about awarding liquidated damages for negligent conduct.

Thank you again for the opportunity to speak on behalf of the physicians of Vermont. The VMS would be happy to review and comment on privacy legislation that comes before the committee. The physicians of Vermont are grateful that this committee has made the privacy of medical records a priority issue.

The CHAIRMAN. Yes. We want to very much. The people who control my life tell me we are running out of time here. [Laughter.]

Ms. MONGAN. Thank you for the opportunity.

Senator LEAHY. YOU have to understand that United States senators are merely constitutional impediments to their staff. [Laughter.] They run everything. [Laughter.]
Senator LEAHY. It is true.
Ms. MONGAN. Thank you.
The CHAIRMAN. You can say that, Pat. I would not dare. [Laughter.]

Senator LEAHY. It is going to cost me, I know. [Laughter.]
The CHAIRMAN. Well, at this point a little off that subject though, Pat and I realize that we have a certain responsibility to Vermont. Senior as we are, Pat is the shadow chairman of the Judiciary Committee, and I am, of course, chairman of the Health, Education, Labor, and Pensions Committee. It gives us an opportunity to get together and to bring staffs up here to listen to Vermont and their problems. It is a great advantage that we share, and we are pledged to doing it as often as we can to make sure that people get the maximum out of our capacities here.

Senator LEAHY. Like the one we had at the Statehouse back here a couple weeks ago, two, 3 weeks ago about Medicaid. We try.
The CHAIRMAN. This is more in your territory. Why don't you—

Senator LEAHY. If I could, Jim, only because I am going to have to slip out for my flight.
The CHAIRMAN. Yes.

Senator LEAHY. Mr. Tofferi, you spoke about individuals having a right to see, and copy and supplement health records and information. I know we talked about doctors and hospital records. What about records on health information that is held by health insurers like your own, should an individual have a right to seek, copy and supplement those records?

Mr. TOFFERI. We do not have a problem with them seeing and copying what we have. We would have an issue with them being able to supplement it. One of the issues the health plan has is different than a medical provider themselves is we do not have the entire medical record for that patient in our possession, and we cannot make decisions that a medical professional could make in certain cases who knows the entire medical record. So our preference is that we be allowed to let them see the record and copy it, but we would also like to have the right—retain the right to require a release from the attending physician or medical professional that we can, in fact, release that information to the patient.

Senator LEAHY. I am wondering if you say you do not have it all, would not that encourage you to ask them to supplement it? You can make the judgment on what weight you want to give to a supplement, but suppose they said, well, now you have got this information about the cancer I had, but you do not have this further information that I have got to have five different checkups all saying I am in total remission, etc., etc. I mean, you might say, okay, fine, we will look at that, but we do not necessarily put a great deal of weight on it because it is such a short period of time or whatever, but would that not be something that might be helpful to you?

Mr. TOFFERI. We only want the minimum amount of information necessary to perform the function. We are not looking to gather additional information. We do not want to be encumbered with storing it, retaining it and all of those kinds of things. Our interest is making sure that we have the minimum but necessary amount of information to actually determine whether it is a covered benefit
and we should pay it or not. If the patient is interested in knowing what we have, we would be happy to give the patient that information with a release from a provider.

Senator LEAHY. You are somewhat unique in the fact you are both an employer and I assume an insurer of some of the same people who work for you. Do you have a firewall to make sure that your employees—or your employers, the supervisors and all, cannot look at an employee's health record unless they are the ones that are supervising the health care plan?

Mr. TOFFERI. Yes, we do. We have a very strict policy in that regard. In fact, we have a separate designated individual in our customer service area who our employees talk to if they have questions about their own benefits. And we are very attentive and very strict about that.

Senator LEAHY. Is that basic industry practice or is that unique to you? It makes a great deal of sense to me; that is why I ask the question.

Mr. TOFFERI. I am only familiar with our own policies, but I would be very surprised if other insurers handled it differently.

Senator LEAHY. You know, we ask these questions and it is easy to bring up the horror stories as I did in my opening statement where it is gone awry where somebody's health records have gone out. The flip of that is we have the best health care system, I believe, in the country. And, Ms. Cramer and Ms. Mongan—am I pronouncing this correctly, Mongan?

Ms. MONGAN. That is correct, Mongan.

Senator LEAHY. When you talk about the patchwork, I am concerned about that. I mean, I know you cross from place to place. You have different laws. My wife who is here is a registered nurse, she is registered both in—she holds licenses both in Vermont and Virginia. You go back and forth on these various things. I look over to Mary Hitchcock or—people move around. Senator Jeffords and I obviously—we have health care here in Vermont, but if something happens when we are in Washington, we have health care there.

If we have a Federal floor, is not it possible to put a Federal floor that evens out almost all of the patchwork but still allows the ability of a State who feels there is an overwhelming compelling purpose to have something stronger or does that create the very problems that you want to avoid?

Ms. CRAMER. I think we are at a time period where as you all know, I am sure, health care is very, very regulated. And we have a variety of type of people who work in a hospital, whether it be the physician, whether it be a technician. So we are constantly training people. And what our goal is so that compliance is as precise as it can be in the environment we work is to have simple, uniform laws. And the difficulty is we do get a lot of turnover. Right now we have shortages in staff. You bring people in for a month, you bring people in for 2 months. We have care that is crossing lines, and it is difficult to always check.

Certainly if Vermont has key priorities, our hospital is routed here, we will know that. Madeleine mentioned the notion of telemedicine. We are all embracing that a little bit more. It brings resources to the State and distributes those resources of those specialists we have here. Sometimes I look at it that we are on this
brink of paralysis sometimes in health care because every direction, we are trying to figure out how five rules sort together. And I am concerned that confidentiality can raise yet another area. And we believe that the penalty should be strong, and so we do not want to have to look at making sure that we have got five states coordinated when our physicians are traveling between New York, and Vermont and Massachusetts.

Senator LEAHY. So your answer to my question is no?
Ms. CRAMER. Correct. [Laughter.]
This is an unusual opportunity.
Senator LEAHY. Hey, that is all right.
Ms. MONGAN. And I could follow up on that no answer with a—

Senator LEAHY. I assume though one thing that may help you is we get better software, better computerization and everything else, but your answer would still be no?
Ms. MONGAN. Yes. And I think it is because in these laws, all of these laws that I have seen at both the Federal and the State level have—there are many small details in them. There are differing time lines, different lengths of time to keep records, different standards for releasing records, the components of what has to be in a release authorizing release of records. I think there is a lot of potential in the details for standards to be different and then for penalties to be imposed. So we are nervous about that, particularly in the light of—I did not get to the penalty section, but in the light of liquidated damages for violation for some these small violations. So we are a little nervous about that.

Senator LEAHY. We will obviously look at all of these. There are a number of competing bills now, competing and complimenting depending how one looks at it. And as I mentioned earlier on, we have that August date. We either do it or it is going to be imposed somehow administratively, and I really would prefer the Congress do it.

I have to leave. We are having some problems in Washington today. They fear snow more than—they fear the threat of snow more than the reality of it, so I have to leave.

Jim, I want to thank you for having this hearing. Obviously you are going to be one that everybody is going to be looking to in the Senate on the issue. It is an extraordinarily important issue. And this underscores something that Senator Jeffords said earlier, we have an advantage in Vermont. We are 600,000 people. We have got the same two senators as California, what do they have now, 35 million, people, and they do not have the opportunity to have hearings like this. You also—in what you are saying though is not just for Vermont. I mean, you become part of the hearing record for the U.S. Senate, and it is in your answers and your statements will be part of that record. And I think we are fortunate that Vermonters take so much time and pay so much attention. I think the rest of the country benefits by it.

Jim, I thank you very much.

The CHAIRMAN. Thanks for dropping by.
Senator LEAHY. Take care.
The CHAIRMAN. Leigh, the Vermont Blue Cross/Blue Shield plan includes a provision explaining its confidentiality practices. I read it. Do you think a patient really understands what it says?

Mr. TOFFERI. I think our members assume that we have a very workable, very tight policy in place. I think they have confidence in our intent in this area. In addition to that, if any member has a particular question, our customer service people are available to go into greater detail. Also, providers who have questions, we have a provider relations staff who can also give them more detailed information on this area. As well as our employer groups. We have a group relations staff who can also provide group administrators with more information on this.

So members may not have detailed information or a detailed understanding, but I think they assume that we have the appropriate standards in effect.

The CHAIRMAN. Ms. Cramer, in your testimony you State that Federal legislation should subject all medical records to the same standards and not provide special disclosures for specific diseases or conditions. How do you suggest we address the concern that mental health records should be at a higher level?

Ms. CRAMER. I think a lot of what we are struggling with are breakdowns in the culture of confidentiality, and I think that there is a lot more that can be done to look at how a medical—mental health record in particular is structured. There have been a lot of informal practices, the description of the notes be included and then not being protected. And so that I think this is an area that is slightly different than the average medical record, but I also think that there are too many primary care providers that need to use this information so that there is not a way to simply have the State Hospital have a set of rules that are stricter and then have individual physicians suddenly exposed to the higher rules. I think we can do a better job in looking at the record, what should be in the record and reconstructing it.

The CHAIRMAN. Thank you. I was the boss of your present boss. Just let him know that I think he does a wonderful job in selecting his fellow attorneys. [Laughter.]

Ms. Mongan, what single issue regarding patient confidentiality is most important to the members of the Vermont Medical Society?

Ms. MONGAN. I think I get a number of calls about just the simple administrative issues from physicians, how long do I have to keep records, how long do I have to keep them after somebody dies and just kind of the—questions about patient's access to records. So I do not know if that is the most important issue, but that is certainly something that physicians would like to have clarified so that there is a clear legal answer, which in Vermont law we do not have answers to a lot of those questions right now. So I think that that is very important.

That is kind of the administrative issue. I think the philosophical point is it is very important to the Medical Society that we protect the confidentiality of patient's records so that that physician-patient relationship can do what it is supposed to do, so that the trust is there between the doctor and the patient that leads to high quality health care.
The CHAIRMAN. Well, thank you, all three of you, for very, very helpful testimony. As I told the other panel, we reserve the right to come back to you with questions. I am sure you will cooperate in providing answers. Deeply appreciate it. Very, very helpful testimony.

Mr. TOFFERI. Thank you.

Ms. MONGAN. Thank you, Senator.

The CHAIRMAN. Our final panelist is Dr. Andrew Gettinger representing Dartmouth-Hitchcock Medical Center in Lebanon, New Hampshire. Dartmouth-Hitchcock is the teaching hospital serving the Dartmouth Medical School, and over 40 percent of its patients are here from Vermont. Thank you for testifying today.

Dr. GETTINGER. Thank you very much.

The CHAIRMAN. Please proceed with your statement.

STATEMENT OF DR. ANDREW GETTINGER, DARTMOUTH-HITCHCOCK MEDICAL CENTER, LEBANON, NH

Dr. GETTINGER. I have submitted written testimony for the record. Rather than reading the written testimony, I would like to give you a sense of who I am as a physician, who we are as an entity in trying to provide health care across State lines.

There has been a lot of discussion already this morning about preemption. Preemption is a complicated issue. We live in two communities where we have fierce protection for individual rights, community rights. It is town meeting season, and we do not give those up without very, very good reason. I think this may be one of those circumstances where there may be good reason.

I provide care to patients who co-exist across State lines. We have defined our medical record as an electronic medical record. Patients receive care concurrently in two states. We cannot distinguish between that care as abiding by rules of one jurisdiction versus another. I was very sensitive to the example of the physician who has offices in three states and patients from all three and the problems that that would entail in regards to preemption.

We believe there is also another aspect to preemption that we would like you to think about; that is that part of our confusion about confidentiality comes from the myriad of different rules and experiences that people try and learn over the course of their lives.

I grew up in New York. I moved to Vermont. I then moved to Connecticut. I moved back to Vermont. I now an a New Hampshire resident, and I do not know what the rules are in each of those entities around the confidentiality of medical information. I am also a teacher, so in preparation for this I went to three sources. I went to my freshman seminar, 12 of the brightest young minds in the United States from 12 different states. They all had different ideas about their personal health information. They did not know. I went to two other sources equally as qualified. I went to the Lime Country General Store on a Saturday morning, [Laughter] and I went to the barber shop in West Lebanon, both locations with people of great intellect. And the diversity of opinion about health care and the confidentiality of medical information was astounding.

Federal preemption also gives us an opportunity to have a level playing field to educate ourselves, to educate our population, our public, and providers need that same level of education. I was very
sensitive to the testimony about having to recruit nurses, respiratory therapists from even outside the United States in order to try and provide health care. We do the same thing.

I am also sensitive to the question you asked about what are the—what are potentially mechanisms that we can put in place to help protect health information. I think there are a variety of mechanisms. I have suggested a couple of them in my written testimony, but I would like to emphasize two of them. The first is that the patient, together with the clinician, physician, author of health information, could in an electronic system identify that information as of higher sensitivity. That would be a joint decision. A flag could then be pushed in the electronic system and that access to that information may result in, first, a warning that this is of a higher level of confidentiality; second, an E-mail back to the author of the information; and third, a request to document why the other clinician needs access to that information.

We have done something similar to that by providing to every employee at Dartmouth-Hitchcock Medical Center access to their own personal health records audit trail, and each clinician who has clinical privileges can as part of their routine day look at who has access to their electronic information and where their paper chart has gone. You know, we have this folksy image of a manila envelope with undecipherable physician encryptions that is absolutely protected from any prying eye, no one including the doctor can re-interpret it. There is no key to re-encyphering this. That has changed. I mean, you heard about $1,500 at some small percent—number of cents per page for health care records that do not fit in a manila envelope anymore. They are mountains.

I have practiced for most of my professional career as an intensivist, that is a physician in the ICUs. In 1 day we can generate more paper than is necessary to provide information. The electronic information systems that we develop I believe can synthesize information out of there, can make that available to other clinicians who need that information when they treat patients.

We believe that patients have strong rights. They are not protected today. You are left to the goodwill of institutions who collect personally identifiable health information to guard that health information. We believe that the computer systems are better than paper systems. We believe that patients can control access to computerized information if we enact guidelines which can serve as a model. We would like to suggest that our affiliate information system is one such model. Physicians, other clinicians regardless of their affiliation with us or not are given access to all the information that we have in the Dartmouth-Hitchcock system about an individual patient, but they are given that information only when the patient consents.

There is an example of that consent form where we try and educate our patients. And the patients always have a right to retract that consent. We believe that is an example of a system that can be endorsed and used more broadly. It enables physicians in the Northeast Kingdom to receive information back about the care that their patients receive at Hitchcock today. Likewise, at Fletcher Allen, similar systems could be in place so that New York physi-


cians and other clinicians can have access to that same information.

I would say that the patchwork of medical information is the disparity between our record keeping systems in all these areas, and I would like you to think about the potential for setting up some uniform standards and an identifier that would enable us in the appropriate settings with appropriate safeguards to aggregate that information in the patient's best interests.

In a short period of time I have covered a variety of different topics.

[The prepared statement of Dr. Gettinger follows:]  

**PREPARED STATEMENT OF ANDREW GETTINGER, M.D.**

Thank you, Mr. Chairman, for the opportunity to testify on this topic of the privacy and confidentiality of personal health information. I am pleased to join my colleagues from Fletcher-Allen Health System, Vermont Blue Cross/Blue Shield, and both the Vermont Medical Society and the Vermont Association of Hospitals and Healthcare Systems. My name is Andrew Gettinger. I am a practicing anesthesiologist, an intensivist (a physician who specializes in critical care medicine) and an active clinical researcher. Additionally, I serve as the Medical Director for Information Systems. In this testimony, I represent the Dartmouth-Hitchcock Medical Center (DHMC) and the Dartmouth-Hitchcock Medical School, which together comprise one of the 126 academic health centers in the United States. The Dartmouth-Hitchcock system provides a full spectrum of care from community-based practices to sophisticated tertiary care. We are the second largest provider of health care to Vermonters, and the largest provider of health care in New Hampshire. For over fifteen years we have worked to develop computerized medical records that assist our clinicians in caring for patients. We are fully committed to addressing the current concerns that accompany the implementation of computers for patient care and research in medicine. That commitment is demonstrated by our adoption of an institutional policy on privacy and confidentiality, fully endorsed by our senior leadership and Boards of Trustees, which outlines strict standards for access to clinical information and mandates serious consequences, including termination, for employees found in violation.

Many in the community have been slow to relinquish the notion of the community-based family doctor whose illegible handwritten notations served as the basis for documenting and recording personal health information. The doctor's manila patient binders were "protected" by this handwritten, illegible scrawl and by the fact that they were located in the doctor's office and not shared with other providers. These patient notes were typically meant for the author as a means to refresh his or her recollection about the patient's personal medical details. Today an individual's health care is provided by a continuum of clinicians for which the timely and accurate conveyance of information is essential. Well-thought-out and well-designed computer systems can facilitate care, not only for the clinician in an academic medical center but also for the clinician in rural settings where patients may be physically distant from referral centers and community physicians isolated from specialists and colleagues. We have developed one such system. Our Affiliate Information System (AIS) provides referring physicians access to the same electronic personal health information as clinicians located at DHMC, but it is the individual patient who controls this access by our requirement of patient consent. The overwhelming majority of our patients grant this consent, understanding that it is in their self-interest to do so.

We believe that there is a clear need for federal legislation to establish the framework for defining and protecting personal health information. The question of federal pre-emption of state law is an important one, especially to our organization. We provide medical care to Vermonters both in Vermont and in New Hampshire. Potentially conflicting state confidentiality statutes could result in circumstances that would be difficult for physicians and health-care providers to comply with; or, worse, impede the flow of critical patient care information. We worry, too, about the development of 50 different standards of health-care confidentiality rather than a single, well-understood structure. It should be as easy for patients to understand the rules regarding personal health information as it is for drivers to navigate our interstate highway system. According to data from the 1990 U.S. census, 34 percent of us reside in states different from the state in which we were born. We have become a
mobile society. Safeguards for privacy and confidentiality need to be structured in a single way, so that patients hear a consistent message and physicians and providers understand clearly their responsibilities. We must be able to educate the population about what privacy standards are and be confident that they understand, whether they are born in Vermont and later move to Washington or grow up in New York and move to New England. Furthermore, health care organizations like DHMC that provide care to residents of two, three or even more states need to have compatible rules to assure consistent protection of confidentiality.

We are in support of legislation to assure privacy and confidentiality of health information. All health care information should be considered sensitive, confidential, and protected so that patients can feel comfortable sharing personal details related to their health. We believe that the answer lies not in segregating data, but in developing mechanisms for protecting the data while allowing access by those who require it to provide the care a patient wants and needs.

We acknowledge that while computer technology can present challenges to the confidentiality of personal health information it can also be implemented in a way that provides security and protection that can not be matched by traditional paper medical records. Two possible safeguards: biometric authentication (that is, having the computer recognize authorized clinicians by their fingerprints, facial identities, or retinal scans) and encryption (that is, scrambling data into bits and pieces that are indecipherable if the key or code is not available) are reaching the stage of development where they are feasible to use.

While it may be premature to rely upon the universal implementation of biometric authentication and encryption for all clinical records, other strategies that are available today can be implemented now. For example, a security structure that allows a variety of access levels based on employee class could be utilized. Clinicians who provide direct care might have the ability to enter, edit and retrieve general patient information. Recognizing that some information, such as HIV status, is considered more sensitive than other information, designated categories of sensitive information might require a higher security level for access. The person requesting access to sensitive data could be required to declare a valid clinical reason for needing to do so. Similarly, other categories of users might only be able to view but not enter or edit information. They too could be required to state the purpose of access to more sensitive information. Both the access and the reasons selected would be reported and monitored by the institution. Certainly, the importance of administrative review and the need to hold all users accountable for their access to information is essential.

Similarly, the physician author of clinical documentation that is particularly sensitive for an individual patient can employ the same approach. Together the patient and physician could choose to put a "restricted" flag on a note that would trigger the procedure described above. The viewer would need to state a reason for requiring access to the note. A message might then be sent back to the author of the note that that it was accessed, by whom, when, and for what reason. The development of such security mechanisms would be facilitated by some national guidelines along these lines, emphasizing security of information, with clear user accountability, rather than segregating data in a way that might have a detrimental impact on patient care.

Individuals and groups concerned about privacy issues have raised great concern in the debate about a national medical identifier. These concerns have been heightened by recent examples of computer hardware and software that have built-in identifiers that exist unbeknownst to the user. Little attention has been paid to the positive outcomes that can result from careful implementation of such a system. Today, personal health information exists in a staggering array of disparate sources that prevent integration and comprehensive medical documentation, all of which would be in the interest of the individual patient. With monitoring, we believe the benefits of a national medical identifier outweigh the potential risks of misuse.

Traditional concepts of research such as randomized controlled clinical trials, case control studies, and patient surveys are overseen by a network of institutional review boards (IRB's). An IRB traditionally functions within an organization with a view to protecting privacy and assuring ethical and sound methods. There is a tension in when trying to maintain the balance between research in the public interest and the individual's right to privacy. Additionally, the distinction between research, which requires IRB oversight, and quality improvement and assurance efforts which do not, is becoming increasingly blurred. We believe that these confidentiality and privacy issues should be addressed by institutions, not through the IRB, but rather through a structure which connects senior leadership, the medical staff, and operational managers so that policy and enforcement can be well-grounded in the daily
operations of the institution. We believe that it is important that these issues be included as the federal legislative process continues.

I am a practicing physician and I care deeply about my patients and their rights. It is important for them, and it is important for me, that they have confidence in the privacy of our interactions and the confidentiality of their health care information. I would like to see federal legislation to assure this protection, while at the same time assuring that complete information is available when needed to those who provide care, and support the provision of care. There is a great opportunity for Congress to legislate such safeguards for privacy and confidentiality that can enhance patient care. We hope that well-publicized but relatively infrequent breaches of confidentiality will not stand in the way of legislation aimed at this greater good. On behalf of our patients in Vermont and New Hampshire, and the physicians and institutions that I represent, we enthusiastically hope and expect that you will complete this legislation in the very near future. We wish you well, and thank you.

The CHAIRMAN. I asked you to come up separately because I am extremely interested in this area. What is going on within the profession or otherwise to figure out how best to address some of these matters?

Dr. GETTINGER. There is a profession, the American Medical Informatics Association, AMIA. I know one of the public speakers is going to represent that. The AMIA has put forward some suggestions. In this month's journal there is actually a coalition, a review article that talks about 10 or 12 principles that might be established. They cite the work in the various bills that have been under consideration, as well as the AMA efforts in this area, as well as Dr. Coop's efforts in these areas as well as the professional societies.

I believe that that kind of information can serve as a very good framework, and I will make that paper available to your staff.

The CHAIRMAN. I would appreciate that, because I am concerned that we have AHCPR, which is one agency that gathers information; and we have the FDA, which is also under our jurisdiction and has a great deal of information. I just imagine a big black box into which all of this information is dumped, and I do not know whether anything comes out that is usable. How do we try to pull this information together? We have additional information flowing in from Europe and Asia.

Dr. GETTINGER. Vermont should take some pride in this. Jack Wennberg, who was one of the leaders in this field, was Commissioner of Medicaid and started his pioneering in variations of health care, how health care that patients receive in Barre are different from that that patients receive in Ludlow. He has continued that work by using the Medicare claims database and has done some very significant work in that area demonstrating differences in health care, the kind of societal differences or regional differences that is not because that we do not all try to do the best we can. We have just not had access to the kind of information that is coming out. What is the best way to take care of Condition A, B or C. I think AHCPR's work is essential in how to get answers to those problems, but in those circumstances you can use the deidentified information. The information does not have to have the patient's name, a medical identifier or any other attributes that really identifies the patients. So I think one can get the public health benefits from quality improvement work with very, very careful application of and rules around patient identifiers and who controls those keys.
The CHAIRMAN. Well, thank you. I am very actively interested and concerned that we are going to end up missing an opportunity to utilize this information much more to our advantage than we currently do.

Dr. GETTINGER. The other point I would like to make just very quickly is that there is consideration of using the IRBs or the CPA chess's, the institutional review boards that regulate research as a venue where confidentiality can be strengthened. I would suggest that this issue is bigger than—by at least an order of magnitude than the IRBs because there needs to be—

The CHAIRMAN. Explain IRBs for the—

Dr. GETTINGER. IRB is the institutional review boards. They tend to be set up around the 126 academic medical centers in the United States. There are some supplemental ones. The FDA has its own IRB. Actually has, I believe, a number of them. And those are the committees that control what researchers can do in their research. They do not today have a clear directive or focus on confidentiality issues per se but are one of the groups that have access to clinical information when it is done under the guise of research.

And so IRBs—my point about IRBs is that they are very important organizations. We need to have them, but I believe this issue goes beyond the IRBs because we have somewhere by an order of magnitude more health care organizations and individual providers, all of whom have information which needs to be protected, by an appropriate framework and standards.

The CHAIRMAN. Well, thank you very much. I will be back in touch with you, because I am very interested in the area you are discussing and how we can maximize our use, at the same time maintaining confidentiality, which is absolutely essential.

Dr. GETTINGER. Thank you for allowing me to testify.

The CHAIRMAN. Thank you.

First on our final panel is Mary Lussier and Ms. Anne Ladue. They are representing the Vermont Occupational Health Nurses Association. The next witness is Dr. Douglas Weir, who has worked since 1983 in IBM Corporation's Occupational Health Services in Essex Junction, Vermont. His medical career includes service at the Veterans Administration Medical Center, Cleveland, Ohio, as well as a residency in internal medicine at the Medical Center Hospital of Vermont. Dr. Weir, nice to have you here. And, finally, I present Dr. Norman S. Ward, who is an associate professor of family practice at the University of Vermont College of Medicine. He is also medical director of Vermont Managed Care.

As a member of the National Health Science Corps early in his medical career, Dr. Ward served in a number of health manpower shortage areas around the country including communities in Mississippi, Georgia, New Hampshire, Maine, and New York State. Since 1995 he has been affiliated with Fletcher Allen Health Care in Burlington. Welcome. Good to have you all here.

And, Ms. Lussier, please proceed.
Ms. LUSSIER. Anne will be reading the statement and I will be answering questions.

Ms. LADUE. Senator Jeffords, my name is Anne Ladue. I am a member of the Vermont chapter of the American Association of Occupational Health Nurses. I am here today to comment on behalf of occupational health nurses and workers across Vermont and the country. AAOHN is the professional association for more than 13,000 occupational health nurses. We provide on the job health care.

AAOHN and I would like to thank you for advancing this very important debate concerning privacy of health information. Enactment of comprehensive Federal legislation is critical. In this age of computers, people are becoming increasingly concerned about their personal health information being used in unintended and potential discriminatory ways.

Today I want to make four points. First, Federal confidentiality legislation should apply to all personally identifiable health information regardless of why the information was generated or where it is collected or stored. Inappropriate disclosure and misuse of health information collected at the work site poses just as serious a threat to the privacy interest of our Nation's workers as does abuse in their treatment, payment, plan administration or research records. Accordingly, effective Federal medical records privacy legislation should cover occupational health records.

Second, human resource managers, line supervisors and others involved in hiring, promotion or work assignment decisions should not have unfettered access to work site medical records. Rather, employers should rely upon the judgment of the health care professionals they retain to generate or maintain occupational health data for determination of an employee's fitness for work. Effective Federal privacy legislation should build barriers designed to prevent management misuse of workers' health records without jeopardizing a company's ability to operate safely and efficiently.

Third, informed authorization for release of medical records should be the cornerstone of any privacy legislation. To be meaningful, the requirements for authorization should apply equally to all types of medical records including those generated or maintained at the work site. That said, a comprehensive Federal privacy bill should include safeguards against coerced authorization for work site records disclosure.

Finally, strong and enforceable civil and criminal penalties for wrongful disclosure or use of protected health information are essential. Moreover, those penalties should apply not only to actual abuses of protected health information but also to inappropriate attempts to access such data. A comprehensive law also should protect individuals who report breaches of the privacy protection from retaliation.
Federal privacy legislation that embraces these four principles should address the privacy concerns that occupational health professionals are forced to grapple with all too often. To illustrate, I would like to relate an incident reported to AAOHN by one of its members. This nurse was threatened with loss of her job when she objected to making employee medical files available to the plant human resources manager so that he could tabulate the frequency and purpose of employee clinic visits for performance appraisal purposes. We need a privacy law that curtails such practices.

The Medical Information Privacy and Security Act includes specific safeguards limiting inappropriate access to employee health information at the work site. The AAOHN is supportive of this level of protection and would like any bill that moves forward to create a similar balance between an employer's legitimate need for employee health information and the employee's right to privacy.

The Vermont chapter and the national office of the AAOHN encourage you and Senator Leahy to work toward the enactment of a bipartisan Federal medical records that is truly comprehensive, recognizing that such a bill must deal effectively with the privacy issue raised by occupational medical records.

We look forward to working with you to achieve this objective. Thank you.

The CHAIRMAN. Dr. Weir.

Dr. WEIR. Thank you, Mr. Chairman, and thank you for the opportunity to present IBM's views on the medical records confidentiality legislation. I would like to begin by condemning your interest and leadership and that of Senator Leahy as well on this issue, and we hope to continue to work with you and your staffs as you develop legislation. We also hope that this Congress will enact such legislation in time for the August 1999 deadline set by the Health Insurance Portability and Accountability Act.

We offer two perspectives on this very important issue: First, as the world's largest information technology company and, second, as an employer with over 150,000 employees in the United States, over 7,000 of which are right here in Vermont.

First I would like to address the role of technology in ensuring privacy and security of our medical records. As a technology company, we see every day the role that information technology plays in improving the delivery of health care, and we believe that such technology can also be used to protect and secure personally identifiable information.

I can imagine a health care system being able to access patient charts and X-rays, attach a physician progress note to an existing electronic chart via voice recognition software, send that file across town or across country to another doctor or hospital. Patients can sit at home and arrange an appointment on line, ask a nurse practitioner about a sick child through an Internet chat session or order prescription refills on the computer. Some of this is available right now.

At the same time, technology offers many forms of security that can appropriately manage access to protected health information, tools such as biometricks and digital signatures that can verify the identity of an individual in a manner that is more fool proof than
a set of questions that ironically enough ask for additional personal information.

Such tools as encryption and other technologies can also enhance the security of health information. To allow our customers to take advantage of the capabilities of the Internet and other technologies, we must create a trusted environment where personal health information is protected. And while we have advocated that in most situations industry self-regulation can address consumer concerns about privacy, for medical records we support Federal legislation to address this important and sensitive category of information. We believe technology can help to implement the goals of this legislation.

Employers regularly play three important and distinct roles relevant to the medical privacy debate. As providers of health benefits and payers of health claims. Self-insured firms like IBM actively design, manage and fund health benefits for their employees; providers of emergency care when necessary and, also, as stewards of a healthy and safe workplace to promote the well-being of their employees.

The employers' role is an active and central one resulting in many positive programs that companies like IBM voluntarily undertake, including various wellness offerings and also disease prevention programs.

It follows then that the employers' benefit plans generate and use much of the health information that flows through the health care system today. In turn, we have a keen interest in the privacy debate.

IBM's own practice aims to maintain a very high standard to ensure confidentiality of information entrusted to it. Our privacy principles relative to employee personnel information are collect, use and retain only personal information that is required for business or legal purposes; provide employees with a means of ensuring that their personal information in IBM personnel records is correct; limit the internal availability of personal information about others to those with a business need to know; and release personal information outside of IBM only with employee approval, except to verify employment or to satisfy legitimate purposes such as investigatory or legal needs.

These principles apply to all personal information but have particular meaning for medical information which, we believe, deserves the greatest degree of protection. Inside IBM access to confidential medical records is limited to the IBM medical staff and it cannot be accessed by management.

To be of benefit, privacy legislation should not tie the hands of employers so severely that they lose the flexibility to perform these necessary functions. Employers need to have access to sufficient information to manage their work force effectively, manage worker’s compensation, comply with the ADA and the FMLA and perform necessary research and analysis that requires medical information. But, clearly, employees need to have assurances that the privacy of their medical records will be protected by their employers.

I will conclude with several points I feel should be addressed in the legislation that Congress enacts on medical records privacy. Legislation should establish a strong Federal framework including
preemption. We cannot encourage development of a sophisticated health care delivery system without uniform national standards for information management.

Privacy and security are closely related, but security standards for health information have already been enacted by Congress in 1996. Final implementing regulations from HHS are expected this year. Federal privacy legislation should not re-legislate these information security standards and therefore create uncertainty, delay implementation and undermine the rule making process.

Federal legislation that sets out uniform standards and responsibilities in this area should strive to create a flexible environment that recognizes the need for a balance of interests. It should also encourage organizations to develop innovative approaches in order to meet their compliance obligations. This flexibility should help spur innovation while at the same time minimizing costs.

Uses of medical information provided by employees to help us as employers make judgments related to ADA, FMLA and other leave of absence programs as a legitimate use and should be accommodated in the legislation.

And while there is certainly a role for strong technology to protect the security of health care information, the legislation should be written in technology neutral terms. This is to allow the flexible use of current technologies while at the same time not discouraging the development of newer alternative technologies in the future. And, finally, privacy legislation should address all types of medical records regardless of the medium in which they are stored.

Once again, I would like to thank you for the opportunity to share IBM's view on Federal medical records confidentiality legislation. And as we have an opportunity to study the legislation recently introduced, we will be very pleased to provide any further comments and answer any questions.

[The prepared statement of Dr. Weir follows:]

PREPARED STATEMENT OF DOUGLAS WEIR, M.D.

Thank you for the opportunity to present IBM's views on medical records confidentiality legislation. My name is Dr. Douglas Weir, and I am a physician who trained in internal medicine at the Medical Center Hospital in Burlington. For the past 15 years I have worked at the IBM facility in Essex Junction where I am currently the Program Manager of IBM Occupational Health Services.

We offer two perspectives on this very important issue: First, as the world's largest information technology company, and second, as an employer with over 150,000 employees in the United States, including over 7,000 in Vermont.

Technology and Medical Records Confidentiality

IBM's Global Healthcare Industry division is a leader in providing new solutions for improving health care systems, patient responsiveness, and communications within far-flung health care operations. We work with our customers—hospitals, providers, insurers, and others—to address their evolving needs and assist them in integrating new technologies into their healthcare systems. I have attached material in the Appendix to this statement that provides more detail on our experience in this industry.

As a technology company, we see every day the role that information technology plays in improving the delivery of healthcare—and we believe that such technology can also be used to protect the privacy and security of personally identifiable information.

As a practicing physician, I can personally imagine a healthcare organization being able to access patient charts and X-rays, attach a physician progress note to an existing electronic chart via voice recognition software, and send that file across town or across country to another doctor or hospital. Patients could sit at home and view the doctor's schedule to arrange an appointment, or ask a nurse practitioner
about a sick child through an Internet chat session. Even using a computer link to determine eligibility and to electronically handle a co-pay while the patient is seeing the doctor. Consumers could even use the Internet or other computer network to buy exercise equipment, or create a diet and exercise plan monitored by an online physician. Some of these are happening even now!

At the same time, technology offers many forms of security controls that can appropriately manage access to protected health information—controls that were not possible or cost-effective in a paper-based environment. Some technology—like biometrics—can authenticate or verify the identity of an individual in a manner that is more foolproof than a set of questions that (ironically) ask for additional personal information. Technology also enables the “de-identification” of information, so that it no longer identifies a specific individual. What that means, in the context of federal privacy legislation, is that it can create information that does not constitute “protected health information” in a manner that is viable for large volumes of healthcare records—something that was not feasible with paper. Encryption and digital signatures are additional components of the technology security portfolio that can be used to enhance the security of health information.

The tools and capabilities I’ve mentioned here will only get better and more sophisticated in the future, given the fast pace of technological advances. In other words, technology can help to implement the goals of this legislation.

Yet this legislation is not directed at these technology solutions, because they are largely addressed under the security standards of the 1996 Health Insurance Portability and Accountability Act’s Administrative Simplification provisions, which are now being implemented by Health and Human Services. Federal medical records confidentiality legislation is directed, instead, at the policies by which organizations will establish who has access to protected information and what the right of individuals are with respect to the uses of the information.

We know as a healthcare information systems company that the only way our customers will be able to take advantage of the capabilities of the Internet and other technologies is if we can create a trusted environment where personal health information is protected. It is to this end that we support the need for and the intent of the legislation, to address this important and sensitive category of information.

An Employer’s Perspective

We all know that most Americans and their families receive health care coverage through their employer—whether that’s a private company or a government. IBM believes that employers play a very constructive role in today’s U.S. health care system and have helped bring about many of the most innovative developments in health care delivery.

Employers regularly play three distinct roles relevant to the medical privacy debate:

• Providers of health benefits and payers of health claims. Self-insured firms like IBM actively design, manage, and fund health benefits for their employees.
• Providers of emergency care when necessary, and
• Stewards of a healthy and safe workplace to promote the well-being of their employees. Employers are responsible for ensuring that a healthy and safe workplace exists at their facilities, and they underpin the nation’s worker compensation system.

Employers’ role is an active and central one, resulting in many positive programs that companies like IBM voluntarily undertake, such as health prevention and wellness programs.

It follows, then, that employers’ benefit plans generate and use much of the health information that flows through the health care system today. In turn, we have a keen interest in the privacy debate.

To be of benefit, privacy legislation should not tie the hands of employers so severely that they lose the flexibility to perform these necessary functions. Employers need to have access to sufficient information to manage their workforce effectively and perform necessary research and analysis that requires medical information. But clearly, employees need to have assurances that the privacy of their medical records will be protected by their employers.

IBM Privacy Policies and Practices

As an information technology company, IBM has maintained a very high standard with respect to ensuring the confidentiality of information entrusted to it. The company’s interest dates back to the 1960s when our privacy policies were formalized due to our desire to respect our individual employees and a general public apprehension about the effects of computers on privacy.

In the 1970s, we conducted a comprehensive review of specific internal guidelines and began management training programs to support compliance with these guidelines.
In the 1980s, we revisited these privacy principles to test their viability given technological and social changes that had occurred and to think through the new challenges presented by these changes.

These are our privacy principles relative to employee personnel information.

• Collect, use and retain only personal information that is required for business or legal reasons.
• Provide employees with a means of ensuring that their personal information in IBM personnel records is correct.
• Limit the internal availability of personal information about others to those with a business need to know.
• Release personal information outside IBM only with employee approval, except to verify employment or to satisfy legitimate purposes, such as investigatory or legal needs.

In the 1990s, and with the explosive growth of the Internet and other networked technologies, we updated our privacy policies by adopting a global online policy for our websites—take a look at our website, at ibm.com. And with the taking effect in October 1998 of the European Union Data Protection Directive, we are again ensuring that our policies meet the requirements of this law.

But we have never lost sight of the fundamental principles, which underlie our employee privacy policies. These principles apply to all personal information but have particular meaning for medical information which, we believe, deserves the greatest degree of protection. Inside IBM, access to confidential medical records is limited to IBM medical staff and department personnel under their immediate supervision. They will disclose information from those records to others within IBM, but only:

• to benefits plan administrators who may review information needed for determining eligibility for benefits
• to others with a need-to-know to evaluate medical recommendations, medical restrictions and accommodations as they relate to the work environment and ability to perform the job
• to legal counsel when medical status or information is at issue or required.

All employees may obtain copies of their records from the IBM medical department. Further, with few exceptions (where required by law or legal process, or where necessary to protect the vital interests of the patient) we require our medical staffs to obtain prior approval of the employee before either disclosing or seeking confidential medical information.

Because we believe that empowered employees with knowledge of their rights is our best assurance that these rules will be followed, we publish our principles and guidelines and periodically remind the 2000+ IBM managers in the U.S. of their responsibilities.

In addition to the information which is contained in our own internally generated medical records, we recognize the need to protect employee medical information associated with our benefits programs. IBM provides a wide array of benefits to our employees, many of which involve treatment for medical conditions—our company provides coverage for over half-million employees, dependents, and retirees. Consistent with our emphasis on employee privacy, we have placed restrictions on our benefits contract administrators on how this information can be used and even what information they will pass on to us. For example, our plan administrators receive only aggregate data derived from the medical records available to the carriers which does not permit linkage of any individual employee with a particular medical condition.

We have imposed these restrictions because we believed it was important to strike the right balance between the needs of the business and the need to protect an employee's privacy. The fact that we have been able to continue to provide our employees a broad array of medical benefits at reasonable costs while operating with these self-imposed restrictions is evidence, we believe, that maintaining high standards of confidentiality need not compromise efficiency.

Federal Legislation Issues

IBM has for years supported federal medical record confidentiality legislation. Personally identifiable medical information is sensitive and deserving of a federal framework for the protection of its privacy and security. We commend Senator Jeffords' and Leahy's interest and leadership on this issue, and hope to continue to work with you and your staffs. We also hope that the new Congress will enact such legislation, in time for the August 1999 deadline set by the Health Insurance Portability and Accountability Act.

Such legislation should include the fundamental “fair information” principles recognized in the privacy debate: notice, access/supplementation, security, and enforcement.
Such legislation should also reflect the following:

Strong Preemption. Legislation should establish a strong federal framework—we cannot encourage development of a sophisticated health care delivery system without national standards for information management.

Health Information Security. Privacy and security are closely related, but as noted previously, security standards for health information have already been enacted by Congress in 1996. Final implementing regulations from HHS are expected this year. Federal privacy legislation should not "re-legislate" these information security standards and therefore create uncertainty, delay implementation and undermine the rulemaking process.

Flexibility. Federal legislation that sets out uniform standards and responsibilities in this area should strive to create a flexible environment, that recognizes the need for balance of interests. It should also encourage organizations to develop innovative approaches in order to meet their compliance obligations. This flexibility should help spur innovation and minimize costs.

Employer's Appropriate Use. We also have a concern over judgments employers must make regarding an employee's ability to perform a job or continue to be eligible for paid or unpaid leave according to company policy. Uses of medical information provided by employees to help us make such judgments is a legitimate use, and should be accommodated in legislation (e.g., there should be no blanket prohibition of the use of information for purposes unrelated to treatment or payment). In fact, proper compliance with laws such as the Americans with Disabilities Act and the Family and Medical Leave Act require that employers make use of personally identifiable medical information in a manner unrelated to treatment or payment.

Technology Neutrality. While there is certainly a role for strong technology to protect the security of health information, legislation should be written in technology-neutral terms. This is to allow the flexible use of current technologies while at the same time not discouraging the development of newer alternative technologies in the future. Finally, privacy legislation should address all types of medical records, regardless of the medium.

Once again, thank you for the opportunity to share IBM's views on federal medical records confidentiality legislation. As we have an opportunity to study the legislation recently introduced, we will be very pleased to provide further input or answer questions.

APPENDIX

IBM & HEALTH CARE INFORMATION TECHNOLOGY

IBM Global Healthcare Industry

As the industry's leading provider of e-business solutions, IBM offers a suite of comprehensive, end-to-end technology solutions to the Healthcare Industry. In the current healthcare marketplace, networking technology now points the way to new methods of improving care systems, patient responsiveness, streamlining cost of operations and improving communications within increasingly far-flung healthcare organizations. IBM's network-enabled commerce, information sharing, management and information technology consulting combined with innovative research offer tangible benefits to healthcare organizations and their constituents.

As healthcare organizations continue to broaden their operations, often including healthcare providers, payers, pharmaceutical and other suppliers in the mix, the need for integrated, secure, scalable networks is leading healthcare organizations towards some form of Internet-based communications.

IBM offers complete solutions suitable for all network configurations: Intranets or internal networks based on local area networks and some form of e-mail and other applications, often built on the Lotus Notes platform; Internet-based programs (Intranets) offering communications and information through websites secured through firewalls and IBM network servers designed for this use; and Extranets which connect Intranets of many organizations through secure Internet channels.

"End-to-end" network computing solutions—from the initial consulting project to implementation and managed operations—address the full spectrum of healthcare organizations' evolving requirements and the need to migrate healthcare systems to new technologies. IBM's 24 development laboratories worldwide work with customers on new technologies such as genomics research to design pharmaceuticals, or continuous speech technology that enables physicians to dictate reports into their computers while reading X-rays or doing a variety of tasks including referrals and other administrative functions.

E-business is the foundation of IBM's network computing offerings providing a flexible, modular approach from a basic foundation level to highly secure, advanced
applications. For those customers who want to create content and establish their presence on the Web, IBM offers solutions including Lotus Notes and Domino™, HTML authoring and HTML templates, TCP/IP and related networking services, web content hosting, e-commerce functions, business recovery services, security, consulting services, and systems management. Our web site development and content hosting services, provide healthcare organizations with a simple-to-operate, engaging set of networked applications offering quality healthcare information and services, personalized to each individual user.

Our Health Data Network provides the framework and solutions whereby payers, providers, government and others share information. The architecture is open and can scale up to support the needs of a growing networked organization. Health Data Network applications enable healthcare organizations to consolidate and access information, automate business processes, minimize redundant data re-entry and extend the useful lives of legacy systems.

Other IBM competitive-advantage healthcare business solutions include data mining to identify marketing trends or ferret out fraudulent claims, etc.; call center technology to handle customer service operations more efficiently at lower costs; mobile computing for enhanced communications with doctors, home healthcare workers, and other healthcare professionals; financial and human resources information systems; year 2000 consulting and software; systems integration and much more.

Professional Service offerings focus on meeting customer requirements which have rapidly increased in scope along with the need to create competitive advantage in a consumer-driven healthcare market environment. Healthcare organizations are looking for sophisticated assessments of Information Systems, HIPAA security readiness, business process evaluation and analysis, reengineering expertise, systems integration, advice on best-of-breed application selection, customization and implementation, ongoing support for such activities, end-user training, and managed operations such as desktop support, data centers, transaction processing, network and other operational business process.

Recognized as the industry's leading provider of electronic business solutions, IBM offers a suite of comprehensive, "end-to-end" technology solutions. IBM's network-enabled e-commerce, information sharing, management and information technology consulting combined with innovative research offer tangible benefits to healthcare organizations and their constituents in improved customer service, enhanced patient care and decreased operating costs.
decisions with my patients. As a medical director, I have been initiated into the complex world of how medical care is paid for. Trying to obtain enough information in a timely enough fashion to approve or disapprove medical services that are or are not benefits within the scope of a patient's insurance contract can be a frustrating experience. As a member of Fletcher Allen's electronic medical information implementation committee, I struggle with the potential wonderful benefits to medical care quality through the use of electronic storage and retrieval of information, balanced by the legitimate concerns of patients for their privacy. As a taxpayer who contributes money to our government, the largest purchaser of health insurance in the country, I am concerned about the high costs of implementing whole new paradigms of information storage and transfer by the health care system.

The medical record is the single most important tool for providing quality health care in 1999. The amazing technologies that can be brought to bear to benefit patients are indeed wonderful, but the medical record is the physician's guide as to when such studies are indicated and appropriate. Since in our current medical system no one practitioner can provide all the care that a patient needs, it is critical that the health care team has ready access to the information that provides the context and background for the next clinical decision. This implies a system with appropriate security safeguards, but one that is highly accessible to the treatment team. We teach our medical students that the key to diagnosis is a thorough and accurate history. Often that history becomes clearer over time such that one small new piece of data makes the whole picture suddenly become clear. The inability to compile information in one usable location significantly handicaps this effort. At the very least, information within an integrated system of care should exist in such a format.

Probably one of the most important aspects of health care today is the use of multiple medications. A patient may see several consultants with different areas of expertise, each of whom may prescribed a medication. Admissions to hospitals for serious complications of medication interactions represent a significant problem. In my own practice, I review the medications that a person is taking on every visit. I instruct the patient to bring every pill bottle in a brown paper bag each time I see them. It is unusual for me not to discover at least one or two discrepancies in what the patient is actually doing with their medications compared to what I believe they are doing. For example, they may be taking both a generic and a brand name preparation of the same drug believing that they are different medications. Our current development of Fletcher Allen's electronic medical information system has placed a high priority on making a shared medication list part of that system to avoid errors and adverse interactions.

Under the proposed legislation, it is conceivable that a patient may choose to restrict access to such a medication list. Let us say, for example, that a patient is on a certain anti-seizure medication. They have blocked their medication list because they are fearful that their employment situation could be threatened if it were known that they had a seizure disorder. They present to the emergency room with an antibiotic known to interact with that medicine
is inadvertently prescribed for a simple infection. The patient may
neglect to inform the emergency room doctor of the fact that they
are on this medication, not realizing the potential for serious drug
interactions. Toxic levels of the seizure medication can result. Now
the patient may return to the emergency room confused and unable
to communicate. The emergency room staff could, I suppose, invoke
the emergency access clause in the legislation to discover the expla-
nation for the symptoms. A preventable complication has occurred.
Where will the liability reside for this untoward event?

Let me speak next about some of the cost implications for the
proposed legislation. There is an enormous cost to the reordering
of tests that may already have been performed when a patient goes
to a new doctor in their own area or more commonly when they are
out of town. Patients often cannot remember the names of tests,
the specific reasons that they were done or the exact results. Sim-
plified systems of accessing this information locally within a pro-
vider network or potentially even nationally in a responsible, se-
cured fashion are important. If State specific statutes create mul-
tiple rules for providers to follow, there will be significant confu-
sion. Fletcher Allen receives some 40 percent of its patients from
New York State. Meeting differing confidentiality statutes for Ver-
mont and New York will prove to be an operational nightmare.

Health care providers are dependent on software vendors to pro-
vide the information systems that they use for patient information.
These systems cost millions of dollars to purchase and install. In-
stallations can take many months. It would be highly desirable if
there were a national standard for the functionality that these sys-
tems need to meet to satisfy privacy requirements. It is conceivable
that a health care system operating in a State with more restric-
tive statutes concerning health information access than a national
norm would find it very difficult to locate a vendor to meet those
specific State needs. In addition, the investments that health care
providers have already made in their current information systems
may be significantly increased by new regulations if those systems
are not compatible with the demands of the new legislation. These
costs will need to be passed on to those paying for medical care.
Strategic decisions made today concerning information technology
capability have implications for years into the future. This speaks
to the need for sufficiently long implementation time line for health
care organizations to attain compliance with any new proposal.

This legislation makes allowance for a patient to sequester parts
of the medical record for transfer. For some people, genetic infor-
mation will be the important category to prohibit access to. For
someone else, it may be a curative cancer operation performed
years before. For another, a notation concerning cosmetic surgery.
This is not to discredit people's individual sensitivities, but rather
to illustrate that the entire record is to be held in a sacred trust.
The logistics of editing a record to the patient's satisfaction prior
to transfer will be daunting.

The legislation also makes reference to eight separate cir-
cumstances where personally identifiable health information can be
released without patient consent. Except for release for emergency
medical treatment, the other seven categories do not involve direct
patient care. Are we to assume that safeguards for confidentiality
within entities such as research, public health, law enforcement and credit card payment systems are more secure than the physicians and nurses providing direct patient care?

In the final analysis, I believe the impetus for this legislation stems partly from heightened concern about patient record confidentiality due to computer related developments. Electronic records can be much more usable as clinical tools than paper records by putting the needed information immediately in the hands of the clinician where the patient is being seen. The ability to document who exactly has opened a computer file and accessed the record can make enforcement of rules prohibiting inappropriate viewing of the record easier. Fletcher Allen has a universal confidentiality statement that all employees must sign stating their understanding that their jobs are at stake for breach of patient privacy.

In closing, I was struck with a sense of irony about the timeliness of the appearance of an article in the Burlington Free Press, you may have seen it, on Saturday in relation to today's meeting. A short article accompanied a picture of Queen Elizabeth documenting the discovery of her medical records in a ditch in Scotland. [Laughter.] Confidentiality is, after all, about people and their imperfections. We are clearly all in this boat together, even the Queen.

I am optimistic that we can, indeed, have a better State of affairs concerning patient privacy in this country. Once again, I wish to thank you for the opportunity to address the committee.

The CHAIRMAN. Thank you. Thank you all. That creates more questions than perhaps I want to deal with, but it is extremely important that we get the best, especially in the area of utilization of technology, in order to sift out some of these problems and get some uniformity.

Do you believe that the bill has sufficient safeguards to allow for uniformity in utilizing technology while insuring both security and the free flow of information? That is a necessity.

Any comments? [Laughter.]

Dr. WARD. Well, if there were a ceiling or if there were at least a floor for what kind of standards, there would need to be a ceiling for the software standards that a vendor would have to meet. Our country is an entrepreneurial society. Those entrepreneurial software manufacturers, for instance, that could exceed the functionality that the national standard puts into place would be at a market advantage, I think. But I think the ceiling should be the important point.

The CHAIRMAN. Dr. Weir.

Dr. WEIR. I agree with what Dr. Ward is saying. I think the most important thing is to have the legislation make clear the importance of confidentiality as a philosophical point. I talked before about not—making it technology neutral because technology changes but the central importance of confidentiality of medical records, and I think that is of all medical records whether they are mental health care, whether they are infectious disease, whatever, is what is important, and it should be unchanging.

The CHAIRMAN. Dr. Gettinger.
Dr. GETTINGER. I believe there is an opportunity with Federal legislation to carefully construct a means to have clinicians, health care providers aggregate data from different sources. Today our problem is that we have bits and pieces of care. Some of your care—some of the documentation of your care lives here in Vermont. Some of the documentation of your care lives in the greater DC. area, and yet any physician who provides care to you either in Vermont or in DC. would like to have all of that information. Standards allowing the aggregation is—are important, and one of the keystones of that standard is a national medical identifier. The problem with a national medical identifier is that everybody is afraid of Big Brother and Big Government. So it is how do you put in place sufficient safeguards and sufficient penalties that will give that kind of number the kind of protection that will make you and me sleep well at night for our own medical information. And I do not think that is a technology specific issue. It is one which will continue on forward.

The CHAIRMAN. It is troubling. A national identifier and availability of all their medical records upsets people. On the other hand, without some sort of system, the maximum protection a patient can have is not available if that information is not locatable and usable.

Dr. WEIR. If I may add something. I think that the concern that a national identifier automatically makes available this myriad of information is incorrect because there are ways to protect that information while it is all clustered under a national identifier.

The CHAIRMAN. Ms. Lussier, will you give me an example of the types of problems the occupational nurses have encountered in treating patients?

Ms. LUSSIER. Some of the problems are that we have collected personal information, medical information along with information that has to do with their occupational job, so we may be asked about some of that which has nothing to do with their job. What we need to do is be able to have the ability just to release information that is needed to say how fit they are to work and that the employer will trust the health care professional that they have the ability to do that. And realize that line managers and supervisors do not have a knowledge to go, and when they go into medical records, to make the right decision what that says and the person's ability to work.

Personally, I have not come in contact with that, but other nurses have come in contact with trying to be forced to release information by the threat of their job, and that we should be able to release only information that is needed. That the employee can sign an authorization saying that they are only to release certain information and that they do not release any other information.

The CHAIRMAN. So you would like to see perhaps some State or Federal laws that would clearly identify or at least set a process for when you can say no to making your medical records available and rely upon the law?

Ms. LUSSIER. Right. And we are protected from having to disclose information; that the health care provider is protected from not being coerced into having to release that information on the threats of their job.
The CHAIRMAN. Ms. Ladue.

Ms. LADUE. I agree with Mary, and many of us nurses work for organizations that are multistate, so it would certainly be to our benefit to have one Federal law to protect so you would not have to know—basically so we would not have to know 50 different laws.

The CHAIRMAN. Well, I want to thank you all for very helpful testimony, although I don't know if it will help us sleep tonight. [Laughter.] But it is making me much more aware of the problems that we have to face. We are still working on the legislation. We would like to be back in touch with you.

Also, I remind everyone here, I told people I would reserve time for questions. So as much as I would like to proceed, I have a promise and I will keep my promise. Thank you for your very excellent testimony.

We have chairs to handle the number of people who have signed up for two-minute statements. We will do it by two panels. The first panel is Andrea Warnke, Vermont ACLU. Laura Ziegler, Phyllis Tarbell, and Vicki Giella. Come up.

We need to breathe for a couple of minutes while the reporter changes the machine and paper.

Because I cannot really connect names with faces, I will proceed from left to right here. Please introduce yourself. Go ahead with your statement.

STATEMENTS OF ANDREA WARNKE, AMERICAN CIVIL LIBERTIES UNION OF VERMONT, MONTPELIER, VT; LAURA ZIEGLER; PHYLLIS TARBELL, DORSET NURSING ASSOCIATION, DORSET, VT; AND VICKI GIELLA, AREA AGENCY ON AGING FOR NORTHEASTERN VERMONT

Ms. WARNKE. Thanks for the opportunity to speak and for your work on these issues. My name is Andrea Warnke. I am the associate director of the American Civil Liberties Union of Vermont. In the interest of leaving time, I am going to abbreviate my comments, but I have a fuller written testimony prepared.

The ACLU obviously is very interested in having a national policy which protects medical records privacy enacted, certainly before the August deadline. The major points of concern to us are that access to patients' medical records should never be given to anyone other than their immediate health care provider without the patients' permission. Use of patient identifiable information should not be permitted unless the patient has granted explicit permission to do so, and the use of unique patient identification numbers, Medicaid or Medicare numbers, or Social Security numbers should be prohibited. Failure to grant such permission should not result in the loss of access to medical care.

Information that you tell your doctor should have the same legal protections as information you tell your lawyer, and law enforcement access to medical records must be limited. Medical records should not become part of a centralized law enforcement database. There must be protective legal standards for access to such records. Law enforcement officers should be able to obtain medical records only after persuading a judge that specific records were necessary for investigating a crime.
State laws which offer greater medical record privacy protection to help their patients must be protected and not preempted. We believe that a Federal standard should set the minimum, not the ceiling.

And, in addition, ACLU is concerned that the use of health care information be restricted even if most personally identifiable information has been removed. And ACLU also believes that after the fact notice should be given in the rare instances where health information has been disclosed to third parties, such as law enforcement agencies, without notice to or consent by the patient. Such a provision should also allow patients to pursue appropriate legal remedies.

Thanks for your time.

[The prepared statement of Ms. Warnke follows:]

PREPARED STATEMENT OF ANDREA WARNKE, ASSOCIATE DIRECTOR, AMERICAN CIVIL LIBERTIES UNION OF VERMONT

The American Civil Liberties Union of Vermont applauds the efforts of our Vermont senators in working to put in place ironclad safeguards to ensure the fundamental privacy rights which all Vermonters expect and deserve with regard to the confidentiality of their medical records.

Privacy protections of medical records have not kept pace with the rapid sweep of technology, although the corresponding industries related to medical data collection have leapt forward with very little oversight and with certainly no comprehensive and consistent national policy protecting the use of such data.

The ACLU believes that we need to immediately enact a national policy protecting medical records privacy which is based on the following principles:

• Access to patients' medical records should never be given to anyone other than their immediate health care provider without the patients' permission. Informed consent should be required before a patient's records are disclosed to third parties except under carefully outlined circumstances. Basic privacy rights can not be violated in the interest of public health and research, which can be conducted without such privacy violations.

• Use of any patient-identifiable information should not be permitted unless the patient has granted explicit permission to do so, and the use of "unique patient identification numbers," Medicaid or Medicare numbers, or Social Security numbers should be prohibited. Failure to grant such permission should not result in the loss of access to medical care. Home health care agencies are now being ordered to be in compliance with federal reporting requirements under OASIS (Outcome and Assessment Information Set) before the end of April. Under OASIS, home health care agencies are being required to electronically submit—for all clients served—patient-specific information including patient name, Medicaid or Medicare number, Social Security numbers, and diagnosis and treatment information. The home health care agencies have been told that failure to provide this information (i.e., if a patient refused permission) would result in prohibiting the patient from receiving treatment. We need a national policy that would prohibit such massive violations of privacy and the punitive withholding of medical care for those who resist.

• Information that you tell your doctor should have the same legal protections as information you tell your lawyer, and law enforcement access to medical records must be limited. Medical records must not become part of a centralized law enforcement database without protective legal standards for access. Law enforcement officers should be able to obtain medical records only after persuading a judge that specific records were necessary for investigating a crimes.

• State laws which offer greater medical record privacy protection to health care patients must be protected, not preempted. Any portion of state laws which provide for a higher standard of privacy protection should take precedence over the national standard, which should set the minimum standard, not the ceiling.

Based on the foregoing statement of principles, ACLU supports the Leahy-Kennedy-Markey bill known as the Medical Information Privacy and Security Act. We would not support other proposed bills until they include, at a minimum, these basic principles.
In addition, ACLU urges that the Medical Information Privacy and Security Act be strengthened by providing for restrictions on the use of health research information even if most personally identifiable information has been removed. ACLU also believes that after-the-fact notice should be provided in the rare instances when health information has been disclosed to third parties such as law enforcement agencies without notice to or consent by the patient. Such a provision should also allow patients to pursue appropriate legal remedies.

We thank you for the opportunity to present our testimony on this critically important issue, and look forward to hearing a response from Senator Jeffords and Senator Leahy on the concerns which we have outlined above.

The CHAIRMAN. Thank you.

Ms. ZIEGLER. I have a stack of paperwork which is unnecessary to say a lot—

The CHAIRMAN. Sure.

Ms. ZIEGLER. The two points I would focus on. I have something which was commissioned by the National Institute of Mental Health from the Mental Health Project in 1978. Part of their model statute included the right to have your records expunged with a court action or sealed. It was at the discretion of the court, but one reason I think it is so important—I should say my background. I was assistant to the director of a mental disability law clinic that served as a protection advocacy office. I also more or less coordinated an advocacy organization that was for and by people who had been locked up in mental institutions. I have a lot of experience with highly confidential information.

In the course of doing discovery I went through massive quantities of this kind of data, and the overwhelming thing that struck me was how pejorative and how often inaccurate this stuff was and the incredible impact it could have on someone’s life, whether through disclosure, shared with providers or with the person’s knowledge that this stuff was on paper. It is kind of like having stuff tattooed on you. Twenty years later what was said about you, which was simply hearsay or which is blatant falsehood and fabrication, somehow has the force of absolute truth because it is in a chart, and the right to put an amendment saying that is not true. Well, I have seen people in court saying that is not true when the chart is cited as what has been going on, and nobody believes them. I am talking about people who have psychiatric labels or histories. So I would strongly recommend that you add this to the repertoire because I think without it, the remedy is extremely inadequate.

I also think that the preemption is not a good idea in terms of you will probably come up with the lowest common denominator for legal rights. I think there is a real need for states to come up with more especially because some states may have model things that are good and innovative, and no one will be able to try them. The State of Michigan granted absolute right of access to records a couple of years ago. They would not have been able to do that if there was Federal statute.

The third thing is that I think that one of the issues that I came across which I do not hear raised, except today it was raised not as let us do something about it but just as part of the landscape, is the illegibility of records. You have no access to something you cannot read. And a staggering amount of what I went through in that discovery took ages just to try and decipher. One of the things that occurred to me is what would happen if it was between health
care providers and how incredibly dangerous. Let us say it was an allergic reaction to a drug or the amount prescribed. There should be something in the statute that requires notations to be in print rather than in cursive. I do not think that would be an unfair burden on anyone. To have signatures have print underneath and to have anything prior to that if it was enacted that you have a right to an explanation and a translation. People have talked about doctors' handwriting and they make a joke out of it, but it is no joke. If you want to see your own records and you cannot read them, and that has certainly happened to me, you never saw them.

I think I have taken up more than 2 minutes.

The CHAIRMAN. All right. We will certainly take your document. Ms. TARBEll. I am Phyllis Tarbell. I am a nurse. I work for the Dorset Nursing Association. We are the smallest home health agency in Vermont, and I am also representing the Vermont Assembly of Home Health Agencies which is composed of the 13 not for profit Medicare certified agencies in the State. We serve between 15 and 20,000 Vermonters each year.

We are very concerned about the new OASIS regulations. Home health agencies are presently mandated to collect and transmit to the State of Vermont and thence to the Health Care Financing Administration OASIS data on every home health patient no matter what the pay source. OASIS records are very personal and sensitive patient identifiable medical information, far more personal than U.S. Census records which are sealed by law for decades. For instance, the OASIS includes all medical diagnoses; all current medications; mental clarity and behavior such as confusion, disorientation, verbal or physical aggression; high risk factors such as obesity, smoking, alcohol or drug dependency; patient's prognosis and even bowel and bladder habits are included along with each patient's name, Social Security Number, their Medicare and Medicaid numbers and their financial status. And all Medicare certified home health agencies in the Nation are required by the new OASIS regs to encode this very private information to transmit it electronically to their State and subsequently to HCFA starting April 26th, 1999, and to electronically update those records every 2 months thereafter at least.

Every patient's right to privacy is being invaded by including those patient's names and identifiers in the documents. And I believe that the Federal government should not have a warehouse of these identifiable, confidential medical records and that they have no right to receive and maintain those records without the patient's informed consent. And when patients become aware that the State and Federal government will be keeping the electronic files of their identifiable records, I think they will have some strong opposition.

When patients refuse to allow the home health agency to transmit their OASIS information, HCFA has decided the agencies must stop providing care to those patients. Agencies will be forced either to abandon patients who refuse to allow transmission of their personal information to HCFA or the agency can risk financial ruin by losing Medicare certification and, therefore, all Medicare revenue.

All Vermont not for profit home health agencies have a mission to serve all in need of home care based on their medical needs, not on their ability to pay or their pay source. We cannot and we will
not abandon our patients if they exercise their right to refuse to provide their OASIS information to HCFA. Agencies that continue to provide care will be violating Medicare conditions of participation and could lose their Medicare certification. It is just unbelievable to me that in these United States patients can be coerced to provide this information or have their coverage denied or their care terminated.

There is a simple solution to this problem. Eliminate the patient names and the identifiers in the encoded and transmitted OASIS records. Agency assigned and protected unique patient numbers already included on each OASIS record would enable investigation of potential patient specific problems by State or Federal officials on a need-to-know basis.

These goals can be accomplished—the goals of OASIS were to authorize—let us see, to require that the home health agency submit information that the Secretary of Health and Human Services considers necessary to develop a reliable case mix system to establish a prospective payment system and to achieve a broad based measurable improvement in quality of care through Federal programs and they can—those goals can be met without including patients' names in that information.

What if anything can we do about this before April 26th? If home health agencies are the only advocates of patients who are not always able to advocate for themselves in every case, we can be viewed as self-serving, i.e., trying to get out of doing OASIS. That is not the problem that I have. Agencies have been cautioned not to present this information to patients in ways that might influence them to refuse permission. In good conscience I could not encourage a patient to consent to this unnecessary invasion of their privacy. Could you?

My colleagues and I will be writing to the Secretary and to HCFA to ask them to eliminate patient names and identifiers from the OASIS before we give HCFA a Pandora's box of patient medical records on April 26th.

Perhaps Congress could persuade them more quickly and effectively than we can possibly hope to. Thank you very much.

The CHAIRMAN. Thank you very much.

Ms. TARBELL. There is a copy of my OASIS document.

[The prepared statement of Ms. Tarbell follows:]

PREPARED STATEMENT OF PHYLLIS TARBELL, RN, BSNA,

Dear Senators and Public, are you aware of the invasion of each individual's right to privacy and the loss of patient confidentiality created by the new home health OASIS requirement under current law. Home health agencies are presently mandated to collect and transmit to the State of VT and thence to the Health Care Financing Administration (HCFA) which oversees Medicare and Medicaid, OASIS data on every home health patient, no matter what the pay source, (except minors and prenatal and postpartum mothers).

OASIS records are very personal and sensitive patient identifiable medical information (see enclosed copy of the questions), far more personal than detailed US Census records which are sealed by law for decades. For instance, the OASIS includes all medical diagnoses, all current medications, mental clarity and behavior, such as confusion, disorientation, verbal or physical aggression, delusional or paranoid behavior. The patient's prognosis, life expectancy, high risk factors such as obesity, smoking, alcohol or drug dependency, even bowel and bladder habits, are all included, along with each patient's name, Soc. Sec. number, Medicare, Medicaid numbers and the patient's financial status. All Medicare certified home health agen-
cies in the nation are required by the new OASIS regulations to encode this very private information and transmit it electronically to their state and subsequently to HCFA, starting April 26, 1999 and to electronically update those records at least every 2 months thereafter.

Every patient's right to privacy is invaded by including patient names and other identifiers in the OASIS document. The federal government should not have a warehouse of patient identifiable, confidential medical files on every home health patient covered by Medicare, Medicaid, private insurance or even private pay. HCFA has no right to receive and maintain personally identifiable OASIS files on patients, particularly those not even covered by Medicare or Medicaid, unless patients give their informed consent for transfer of that information. When patients become aware that the state and federal governments will be maintaining electronic files of their personally identifiable, private and confidential medical records, I think there will be very strong opposition.

If patients refuse to allow the home health agency to transmit their OASIS information, then HCFA has decided that agencies must stop providing care to those patients. Agencies will be forced either to abandon patients who refuse to allow transmission of their personal information to HCFA, or risk financial ruin by losing Medicare certification and all Medicare revenue.

All Vermont not for profit home health agencies have a Mission to serve all in need of home care, based on their medical needs not on ability to pay or pay source. We cannot and will not abandon our patients if they exercise their right to refuse to provide their OASIS information to HCFA. Agencies that continue to provide care to those patients will be violating Medicare Conditions of Participation and could lose their Medicare certification. It is unbelievable that in these United States patients can be coerced to provide this information, or have:

1. Medicare/Medicaid coverage denied
2. Medically necessary care terminated even if they are not covered by MC/MA or choose to pay privately.

There is a simple solution to this problem; eliminate patient names and other identifiers from all encoded and transmitted OASIS records. Agency assigned and protected unique patient numbers already included on each OASIS patient record would enable investigation of potential patient specific problems by state or federal officials on a need to know basis. The Federal Register 1/25/99, states (quote): "Summary: "Section 4602(e) The Balanced Budget Act authorized the Secretary of the Dept of Health and Human Service) to require that home health agencies submit any information the Secretary considers necessary to develop a reliable case mix system. . . . to establish a prospective payment system (PPS) for home health agencies and to achieve a broad based measurable improvement in quality care furnished through Federal programs." Did Congress really mean "any Information the Secretary considers necessary"? That is far too much freedom to decide what might or might not be necessary to achieve any goal.

These goals can be accomplished without patient names and other personal identifiers encoded and submitted in the OASIS records. There is no justification or need to include patient identification routinely for anyone except direct medical care providers to see. Medicare surveyors that regularly inspect home health agencies have always had access to every home health agency patient's medical record during surveys and when problems or complaints arise. But never before have identifiable patient medical records been copied, removed from the agency, and constantly updated and maintained by the state and federal governments for government staff to access and retrieve.

The nursing home industry has already been forced to provide similar personal, private medical information (MDS) on every Medicare or Medicaid patient they serve. Home health is now to send information (OASIS) on every patient regardless of payment source (except minors and prenatal and postpartum mothers). There is, I understand, a progressive plan for more and more health care providers, including hospitals, to send HCFA their patient information, until eventually the federal government will have personally identifiable medical files on each and every one of us. It is none of their business and it is wrong.

What, if anything, can we do about it before April 26, 1999? If home health agencies are the only advocates of patients who are not always able to advocate for themselves, we will be viewed as self serving, ie "trying to get out of doing OASIS". Agencies have been cautioned not to present this information to patients in ways that might influence them to refuse permission. In good conscience I could not encourage a patient to consent to this unnecessary invasion of their privacy. Could you?

I and my colleagues will be writing to ask the Secretary and HCFA to ask them to eliminate patient names and identifiers from OASIS, before we give HCFA a Pandora's box of patient medical records on April 26.
But perhaps Congress could persuade them more quickly and effectively than we can possibly hope to.

Thank you.
OASIS Items: all those starting with "mo---"
**COMPREHENSIVE START OF CARE ASSESSMENT**

(Also used for Resumption of Care Following Inpatient Stay)

**[MD7799]** From which of the following Inpatient Facilities was the patient discharged during the past 14 days? (Mark all that apply.)

- Hospital
- Rehabilitation facility
- Nursing home
- Other (specify)
- Patient was not discharged from an Inpatient facility [If NA, go to MD7999]

**[MD7999]** Inpatient Diagnosis Date (must be recent)

**[MD7999]** Inpatient Diagnoses and ICD code categories (three digits required; five digits optional) for only those conditions treated during an Inpatient facility stay within the last 14 days (no surgical or V-codes):

<table>
<thead>
<tr>
<th>Inpatient Facility Diagnosis</th>
<th>ICD</th>
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</thead>
<tbody>
<tr>
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</table>

**[UM2296]** Medical or Treatment Regimen Change Within Past 14 Days: Has the patient experienced a change in medical or treatment regimen (e.g., medication, treatment, or service change due to new or additional diagnosis, etc.) within the last 14 days?

- No (If No, go to UM2297)
- Yes

**[UM2296]** List the patient's medical diagnoses and ICD code categories (three digits required; five digits optional) for those conditions requiring changed medical or treatment regimen (no surgical or V-codes):

<table>
<thead>
<tr>
<th>Changed Medical Reason Diagnoses</th>
<th>ICD</th>
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</thead>
<tbody>
<tr>
<td></td>
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</table>

**[UM2296]** Conditions Prior to Medical or Treatment Regimen Change or Inpatient Stay Within Past 14 Days: If the patient experienced an Inpatient facility discharge or change in medical or treatment regimen within the past 14 days, indicate any conditions which existed prior to the Inpatient stay or change in medical or treatment regimen. (Mark all that apply.)

- Urinary incontinence
- Indwelling/urinary catheter
- Intractable pain
- Impaired decision making
- Disruptive or socially inappropriate behavior
- Memory loss to the extent that supervision required
- None of the above
- NA: No Inpatient facility discharge and no change in medical or treatment regimen in past 14 days

**[UM2299]** Diagnoses and Severity Indicators: List each medical diagnosis or problem for which the patient is receiving home care and ICD code category (three digits required; five digits optional, no surgical or V-codes) and rate them using the following severity index. (Choose one value that represents the most severe rating appropriate for each diagnosis.)

0 - Asymptomatic, no treatment needed at this time
1 - Symptoms well controlled with current therapy
2 - Symptoms controlled with difficulty, affecting daily functioning, patient needs ongoing monitoring
3 - Symptoms poorly controlled, patient needs frequent adjustment in treatment and dose monitoring
4 - Symptoms poorly controlled, history of rehospitalizations

**[UM2299]** Primary Diagnoses

<table>
<thead>
<tr>
<th>ICD</th>
<th>Severity Rating</th>
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<td>0 0 0 0 0 0 0</td>
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</table>

**[UM2299]** Other Diagnoses

<table>
<thead>
<tr>
<th>ICD</th>
<th>Severity Rating</th>
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<td></td>
<td>0 0 0 0 0 0 0</td>
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</table>

**Key:**

- More than one response allowed
- Only one response allowed

## COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

### THERAPIES the patient receives at home (Mark all that apply)
- 1. Intravenous or Infusion Therapy (excluding TPN)
- 2. Parenteral nutrition (TPN or lipid)
- 3. Enteral nutrition (enterostomy, gastrostomy, or any other artificial entry into the alimentary canal)
- 4. None of the above

### Overall Prognosis: BEST description of patient's overall prognosis for recovery from the episode of illness
- 0. Poor: little or no recovery is expected and/or further decline is imminent
- 1. Good: partial to full recovery is expected
- 2. Unknown

### Rehabilitation Prognosis: BEST description of patient's prognosis for functional status
- 0. Declined: minimal improvement in functional status is expected, decline is possible
- 1. Good: marked improvement in functional status is expected
- 2. Unknown

### Life Expectancy: (Physician documentation is not required)
- 0. Life expectancy is greater than 9 months
- 1. Life expectancy is 9 months or fewer

#### ADVANCE DIRECTIVES:
- Mark all that apply
- Patient does not have Advance Directives
- Patient has Advance Directives
- Written information provided to Patient / caregiver
- Copy provided for clinical record

### HIGH RISK FACTORS characterizing this patient
- Mark all that apply
- Heart smoking
- Obesity
- Alcohol dependence
- Drug dependence
- None of the above

### HEALTH SCREENING
- Patient receives preventive and/or periodic health screenings
- Patient referred for appropriate preventive/periodic health screenings

#### Guidelines (individual risk factors & diagnosis dictate actual follow-up)
- 1. Age 50 year or older after normal tests.
- 2. Age 50 year or older after abnormal tests.
- 3. Mammography: 0 year or age 50
- 4. Cervical / Rectal 0 year or age 50
- 5. Immunizations for adults (flu shot annually, tetanus q/5y, pneumococcal once in a lifetime. Meningococcus q y
- 6. Physical exam 0 year or age 50

#### LIVING ARRANGEMENTS

#### Current Residence
- 0. Patient's owned or rented residence (house, apartment, or mobile home owned or rented by patient
- 1. Family member's residence
- 2. Boarding home or rented room
- 3. Board and care or assisted living facility
- 4. Other (specify)

#### Structural Barriers in the patient's environment
- Mark all that apply
- No
- Stairs inside home which must be used by the patient (e.g., to get to kitchen, sleeping, eating areas)
- Stairs inside home which are used optionally (e.g., to get to laundry facilities)
- Narrow or obstructed doorways

#### Safety Hazards found in the patient's current place of residence
- Inadequate floor, cool, or windows
- Inadequate lighting
- Unsafe heating appliance
- Inadequate heating
- Inadequate cooking
- Lack of fire safety devices
- Unsafe floor coverings
- Inadequate stair railings
- Improperly stored hazardous materials
- Lead-based paint
- Other (specify)
## COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

### Safety Plan (Mark all that apply):
- Outdoor natural disaster plan, including system for priority status.
- Emergency procedures for medical emergencies, and fire plan.
- Instructed in fall prevention measures.
- Referral to physical therapy.
- Patient/Caregivers instructed in managing risks within environment.

### Patient Lives With: (Mark all that apply, select one)
- Lives alone
- With spouse or significant other
- With other family member
- With a friend
- With paid help (other than home care agency staff)
- With other than above

### Assistive Person(s) Other Than Home Care Agency Staff:
- Relative, friends, or neighbors living outside the home
- Person residing in the home (EXCLUDING paid help)
- Paid help
- None of the above

### Primary Caregiver having legal responsibility for providing or managing the patient's care, providing the most frequent assistance, etc. (other than home care agency staff):
- No one person
- Spouse or significant other
- Daughter or son
- Other family member
- Friend or neighbor or community or church member
- Paid help
- Unknown

### How Often does the patient receive assistance from the primary caregiver?
- Several times during the day and night
- Several times during the day
- Once daily
- Three or more times per week
- One to two times per week
- Less often than weekly
- Unknown

### Type of Primary Caregiver Assistance: (Mark all that apply)
- ADL assistance (e.g., bathing, dressing, toileting, Bowler, feeding)
- Medical assistance (e.g., meals, meds, housekeeping, laundry, cleaning)
- Environmental support (housmg, home maintenance)
- Psychosocial support (depression, loneliness, recreation)
- Advocates or facilitates patient's participation in appropriate medical care
- Financial agent, power of attorney, conservator of property, or medical power of attorney
- Unknown

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**Key:**
- D - more than one response allowed
- O - only one response allowed

SOC Assessment Incorporating OASIS-B1 (1996)
## COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

### SENSORY STATUS

**EYES**
- [ ] Photosensitivity
- [ ] Photophobia
- [ ] Glaucoma
- [ ] Cataract R/L
- [ ] Blurred Vision R/L
- [ ] Double Vision R/L
- [ ] Other (specify) __________________________
- [ ] Not assessed

**SENSORY STATUS**
- [ ] Normal vision: sees adequately in most situations; can see medication labels, newspapers
- [ ] Partially impaired: cannot see medication labels or newspapers, but can see objects in field of vision
- [ ] Severely impaired: cannot locate objects without hearing or touching them; is patient nonresponsive

**EARS**
- [ ] Normal hearing: able to hear and understand questions and instructions
- [ ] Hearing loss: needs amplification
- [ ] Hearing aid: needs amplification
- [ ] Other (specify) __________________________
- [ ] Not assessed

**ORAL**
- [ ] Speech and Oral (Verbal) Expression of Language (in patient's own language):
  - [ ] Expresses complex ideas, feelings, and needs clearly, completely, and easily in all situations
  - [ ] Expresses simple ideas or needs with moderate difficulty
  - [ ] Expresses ideas or needs with severe difficulty
  - [ ] Unable to express basic needs even with maximal assistance or assistance needed is not consistent or unresponsive,
    or I/O speech is incoherent or unintelligible
- [ ] Other (specify) __________________________
- [ ] Not assessed

**HEMATOPOETIC ASSESSMENT**
- [ ] Excessive bleeding/bruising
- [ ] Not assessed

## DERMAL ASSESSMENT
- [ ] Rash
- [ ] Scaling
- [ ] Poor Turgor
- [ ] Not assessed

---

**HEMATOPOETIC ASSESSMENT**
- [ ] Excessive bleeding/bruising
- [ ] No
- [ ] Yes

**DERMILL ASSESSMENT**
- [ ] Rash
- [ ] Scaling
- [ ] Poor Turgor
- [ ] Not assessed

---

Not more than one response allowed
SOC Assessment Incorporating OASIS-8.1 (1099)

O = only one response allowed

---

5
**COMPREHENSIVE START OF CARE ASSESSMENT**

(Also used for Resumption of Care Following Inpatient Stay)

(M6448) Does the patient have a Skin Lesion or an Open Wound? This excludes "STOMIES."

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
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<td>O 0</td>
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<tr>
<td>O 1</td>
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</table>

- If skin lesion or open wound is other than a pressure or stasis ulcer, or a surgical wound, describe an outcome planner.

(M6449) Does this patient have a Pressure Ulcer?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
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<td>O 0</td>
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<tr>
<td>O 1</td>
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</tbody>
</table>

(M6450) Current Number of Pressure Ulcers at Each Stage: (Circle one response for each stage)

<table>
<thead>
<tr>
<th>Stage</th>
<th>Number of Pressure Ulcers</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0 1 2 3 4 or more</td>
</tr>
<tr>
<td>2</td>
<td>0 1 2 3 4 or more</td>
</tr>
<tr>
<td>3</td>
<td>0 1 2 3 4 or more</td>
</tr>
<tr>
<td>4</td>
<td>0 1 2 3 4 or more</td>
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</table>

(M6444) Stage of Most Problematic (Observable) Pressure Ulcer:

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<thead>
<tr>
<th></th>
<th>Stage 1</th>
<th>Stage 2</th>
<th>Stage 3</th>
<th>Stage 4</th>
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<tbody>
<tr>
<td>O 0</td>
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(M6451) Status of Most Problematic (Observable) Stasis Ulcer:

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4 or more</th>
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<td>O 0</td>
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<td>O 3</td>
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(M6445) Does this patient have a Stasis Ulcer?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
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<tbody>
<tr>
<td>O 0</td>
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<tr>
<td>O 1</td>
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</table>

(M6447) Current Number of Observable Stasis Ulcers:

<table>
<thead>
<tr>
<th></th>
<th>Zero</th>
<th>One</th>
<th>Two</th>
<th>Three</th>
<th>Four or more</th>
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<tbody>
<tr>
<td>O 0</td>
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<td>O 1</td>
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<td>O 3</td>
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</tbody>
</table>

(M6446) Does this patient have at least one Stasis Ulcer that Cannot be Observed due to the presence of a non-removable dressing?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>O 0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O 1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(M6448) Status of Most Problematic (Observable) Surgical Wound:

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4 or more</th>
</tr>
</thead>
<tbody>
<tr>
<td>O 0</td>
<td></td>
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<td>O 1</td>
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<tr>
<td>O 2</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>O 3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(M6452) Does this patient have a Surgical Wound?

<table>
<thead>
<tr>
<th></th>
<th>No</th>
<th>Yes</th>
</tr>
</thead>
<tbody>
<tr>
<td>O 0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O 1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(M6449) Current Number of Observable Surgical Wounds:

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4 or more</th>
</tr>
</thead>
<tbody>
<tr>
<td>O 0</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>O 1</td>
<td></td>
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<td></td>
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<tr>
<td>O 2</td>
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<td></td>
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<tr>
<td>O 3</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

(M6447) Does this patient have at least one Surgical Wound that Cannot be Observed due to the presence of a non-removable dressing?

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>O 0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>O 1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(SOC Assessment Incorporating OASIS-81 (1998))
COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

1. NUTRITIONAL STATUS (Mark all that apply): If any risk factors are identified, appropriate follow-up is needed.
   - Special Diet: ____________________________
   - Open wound
   - Poor nutrition
   - Oral health problems
   - Food preferences
   - Other (specify) ____________________________

2. CARDIOVASCULAR ASSESSMENT (Mark all that apply):
   - Hemodynamic status
   - Fatigue
   - Other (specify) ____________________________
   - Peripheral vascular disease
   - Other (specify) ____________________________

3. RESPIRATORY ASSESSMENT (Mark all that apply):
   - Cough
   - Difficulty breathing in recumbent position
   - Other (specify) ____________________________
   - Not assessed

4. PSYCHOSOCIAL ASSESSMENT (Mark all that apply):
   - Home environment
   - Living situation
   - Other (specify) ____________________________
   - Other (specify) ____________________________

5. PATIENT TREATMENT (Mark all that apply):
   - Oxygen administration
   - Ventilator (cotton-linility or at night)
   - Cardiac pacemaker
   - Other (specify) ____________________________

6. RESIDENT ASSESSMENTS (Mark all that apply):
   - Sensory impairment
   - Hearing impairment
   - Other (specify) ____________________________

7. GASTROINTESTINAL ASSESSMENT (Mark all that apply):
   - Constipation
   - Other (specify) ____________________________

8. BOWEL INCONTINENCE (Specify)
   - Very rarely or never has bowel incontinence
   - More often than once daily
   - Other (specify) ____________________________

9. URINARY INCONTINENCE (Specify)
   - No incontinence or catheter (includes urinary drainage or Foley catheter, if applicable)
   - Other (specify) ____________________________

10. OXYGEN REQUIREMENTS (Specify)
    - Patient does not have an oxygen need related to an institutional stay or it is not necessary change in medical or treatment regimen.
    - Other (specify) ____________________________
# COMPREHENSIVE START OF CARE ASSESSMENT

(Also used for Resumption of Care Following Inpatient Stay)

<table>
<thead>
<tr>
<th>1. NEURO/EMOTIONAL/BEHAVIORAL ASSESSMENT (Mark all that apply.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>□ Frequent or severe headaches □ Slurred speech □ Numbness □ Tremors □ Unsteady gait</td>
</tr>
<tr>
<td>□ Other (specify) □ Drowsy problems □ Not assessed</td>
</tr>
</tbody>
</table>

(65564) Cognitive Functioning: (Patient's current level of alertness, orientation, comprehension, concentration, and immediate memory for simple commands.)

- 0 - Alert, oriented, able to focus and shift attention, comprehends and recalls task directions independently.
- 1 - Requires prompting (talking, repeating, reminding) only under stressful or unfamiliar conditions.
- 2 - Requires assistance in tasks and some direction in specific situations (e.g., on all tasks involving shifting of attention), or consistently requires low attention environment due to distractibility.
- 3 - Requires considerable assistance in routine situations. Is not alert and oriented or is unable to shift attention and recall directions more than half the time.
- 4 - Totally dependent due to disturbances such as constant disorientation, coma, persistent vegetative state, or dementia.

(65679) When Confused (Reported or Observed):

- 0 - Never
- 1 - In new or complete situations only
- 2 - On awakening or at night only
- 3 - During sleep or evening, but not constantly
- 4 - Constantly
- NA - Patient nonresponsive

(65680) Patient Behaviors (Reported or Observed): (Mark all that apply.)

- 0 - None of the above feelings observed or reported
- 1 - Depressed mood (e.g., feeling sad, lonely)
- 2 - Sense of failure or self-reproach
- 3 - Haplessness
- 4 - Recurrent thoughts of death
- 5 - Thoughts of suicide
- 6 - None of the above feelings observed or reported

(65689) When Anxious (Reported or Observed):

- 0 - None of the time
- 1 - Less than half the time
- 2 - Daily, but not constantly
- 3 - All of the time
- NA - Patient nonresponsive

(65690) Patient Behaviors (Reported or Observed): (Mark all that apply.)

- 0 - None of the above behaviors observed or reported
- 1 - Intolerance, lack of concentration
- 2 - Diminished interest in most activities
- 3 - Sleep disturbances
- 4 - Recent change in appetite or weight
- 5 - Agitation
- 6 - A suicide attempt
- 7 - None of the above behaviors observed or reported

(65660) Behavior Demonstrated at Least Once a Week (Reported or Observed): (Mark all that apply.)

- 0 - Memory deficit, failure to recognize familiar persons/objects, inability to recall events of past 24 hours, significant memory loss so that supervision is required
- 1 - Impaired decision-making, failure to perform usual ADLs or IADLs, inability to appropriately stop activities, jeopardizes safety through actions
- 2 - Verbal disruption, yelling, threatening, excessive profanity, sexual references, etc.
- 3 - Physical aggression, aggressive or combative to self and others (e.g., hits self, throws objects, punches, dangerous maneuvers with objects or other objects)
- 4 - Disorganized, infantile, or socially inappropriate behavior (excludes verbal actions)
- 5 - Delusional, hallucinatory, or paranoid behaviors
- 6 - None of the above behaviors demonstrated

(65659) Frequency of Behaviors Problems (Reported or Observed) (e.g., wandering episodes, self abuse, verbal disruption, physical aggression, etc.)

- 0 - Never
- 1 - Less than once a month
- 2 - Once a month
- 3 - Several times a month
- 4 - Several times a week
- 5 - At least daily

(65661) Is this patient currently Psychiatric Nursing Services at home provided by a qualified psychiatric nurse?

- 0 - No
- 1 - Yes

# MUSCULOSKELETAL ASSESSMENT (Mark all that apply.)

- 0 - Weakness
- 1 - Decreased ROM (specify) [specify]
- 2 - Swollen, painful joints (specify) [specify]
- 3 - Gait disturbance
- 4 - Other (specify)
- 5 - Dorsal problems
- 6 - Not assessed

Key: X = more than one response allowed
O = only one response allowed
SOC Assessment Incorporating OASIS-B1 (1999)
COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

LIFE SYSTEM PROFILE: For the "prior" column, mark the box that corresponds to the patient's condition 1-4 days prior to start of (or resumption of) care. In all cases, record what the patient was able to do.

**Grooming**: Ability to tend to personal hygiene needs (e.g., washing face and hands, hair care, shaving or make up, teeth or denture care, fingernail care)

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>0</td>
</tr>
<tr>
<td>O</td>
<td>1</td>
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<tr>
<td>O</td>
<td>2</td>
</tr>
<tr>
<td>O</td>
<td>3</td>
</tr>
<tr>
<td>O</td>
<td>UK</td>
</tr>
</tbody>
</table>

**Ability to Dress Upper Body (with or without dressing aids) including undergarments, pullers, front opening shirts and knits, managing button, buttons, and zippers**

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
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<tbody>
<tr>
<td>O</td>
<td>0</td>
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<td>O</td>
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<td>O</td>
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<td>O</td>
<td>3</td>
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<tr>
<td>O</td>
<td>UK</td>
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</table>

**Ability to Dress Lower Body (with or without dressing aids) including undergarments, socks, socks or shoes, shoes.**

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
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<tbody>
<tr>
<td>O</td>
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<td>O</td>
<td>3</td>
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<tr>
<td>O</td>
<td>UK</td>
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</table>

**Bathing**: Ability to wash entire body. Excludes grooming (washing face and hands only).

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
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<tbody>
<tr>
<td>O</td>
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<td>O</td>
<td>4</td>
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<tr>
<td>O</td>
<td>5</td>
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<tr>
<td>O</td>
<td>UK</td>
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</tbody>
</table>

**Toileting**: Ability to get to and from the toilet or bedside commode.

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
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<tbody>
<tr>
<td>O</td>
<td>0</td>
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<tr>
<td>O</td>
<td>1</td>
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<tr>
<td>O</td>
<td>2</td>
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<td>3</td>
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<tr>
<td>O</td>
<td>4</td>
</tr>
<tr>
<td>O</td>
<td>UK</td>
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</tbody>
</table>

Key: D = one response allowed  O = only one response allowed  EOC Assessment Incorporating OASIS-II (1998)
COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

Transporting:
Ability to move from bed to chair, on and off bed or commode, into and out of tub or shower, and ability to turn and position self is

dependent on patient's condition.

Prior
Current
0 0 0 - Able to independently transfer.
0 0 1 - Transfers with minimal human assistance or with use of an adaptive device.
0 0 2 - Unable to transfer self but is able to bear weight and pivot during the transfer process.
0 0 3 - Unable to transfer self and is unable to bear weight or pivot when transferred by another person.
0 0 4 - Bedfast, unable to transfer but is able to turn and position self in bed.
0 0 5 - Bedfast, unable to transfer and is unable to turn and position self.
0 0 6 - Unable
0 0 7 - Unknown

Ambulation/ locomotion:
Ability to safely walk, once in a standing position, or use a wheelchair, once in a seated position, on a variety of surfaces.

Prior
Current
0 0 0 - Able to independently walk on even and uneven surfaces and climb stairs with or without railings (i.e., no need for human assistance or adaptive device).
0 0 1 - Requires use of a device (e.g., cane, walker) to walk alone or requires human supervision or assistance to negotiate stairs or steps or uneven surfaces.
0 0 2 - Able to walk only with the supervision or assistance of another person at all times.
0 0 3 - Can transfer, unable to ambulate but is able to walk self independently.
0 0 4 - Can transfer, unable to ambulate and is unable to walk self.
0 0 5 - Bedfast, unable to ambulate or be up in a chair.
0 0 6 - Unknown

Feeding or Eating:
Ability to feed self meals and snacks. Note: This refers only to the process of eating, chewing, and swallowing, not preparing the food to be eaten.

Prior
Current
0 0 0 - Able to independently feed self.
0 0 1 - Able to feed self independently but requires:
(a) meal set-up; OR
(b) Intermediary assistance or supervision from another person; OR
(c) a liquid, pureed, or ground meal diet.
0 0 2 - Unable to feed self and must be assisted or supervised throughout the mealtime.
0 0 3 - Able to take in nutrients orally and receives supplemental nutrients through a nasogastric tube or gastrostomy.
0 0 4 - Unable to take in nutrients orally and is fed nutrients through a nasogastric tube or gastrostomy.
0 0 5 - Unable to take in nutrients orally or by tube feeding.

Prior
Current
0 0 6 - Unable to feed
0 0 7 - Unknown

Planning and Preparing Light Meals (e.g., cereal, sandwich) or meal delivered meals:

Prior
Current
0 0 0 - (a) Able to independently plan and prepare all light meals for self or receive delivered meals; OR
(b) Physically, cognitively, and mentally able to prepare light meals on a regular basis but has not routinely performed light meal preparation in the past (i.e., prior to this home care admission).
0 0 1 - Unable to prepare light meals on a regular basis due to physical, cognitive, or mental limitations.
0 0 2 - Unable to prepare any light meals or refused any delivered meals.
0 0 3 - Unable to prepare any light meals or low fat meals due to physical, cognitive, or mental limitations.
0 0 4 - Unable to prepare any light meals or refused any delivered meals.
0 0 5 - Unable to prepare any light meals or refused any delivered meals.
0 0 6 - Unable to prepare any light meals or refused any delivered meals.
0 0 7 - Unknown

Transportation:
Physical and mental ability to safely use a car, taxi, or public transportation (bus, train, subway).

Prior
Current
0 0 0 - Able to independently drive a regular or adapted car. OR uses a regular or handicap-accessible public bus.
0 0 1 - Able to ride in a car only when driven by another person. OR uses a bus or handicap van only when assisted or accompanied by another person.
0 0 2 - Unable to ride in a car, taxi, bus, or van, and requires transportation by ambulance.
0 0 3 - Unable to ride in a car, taxi, bus, or van, and requires transportation by ambulance.
0 0 4 - Unable to ride in a car, taxi, bus, or van, and requires transportation by ambulance.
0 0 5 - Unable to ride in a car, taxi, bus, or van, and requires transportation by ambulance.
0 0 6 - Unable to ride in a car, taxi, bus, or van, and requires transportation by ambulance.
0 0 7 - Unknown

Laundry:
Ability to do own laundry - to carry laundry to and from washing machine, to use washer and dryer, to wash small items by hand.

Prior
Current
0 0 0 - (a) Able to independently take care of all laundry tasks; OR
(b) Physically, cognitively, and mentally able to do laundry and access facilities, but has not routinely performed laundry tasks in the past (i.e., prior to this home care admission).
0 0 1 - Able to do only light laundry, such as hand wash or light washer loads. Due to physical, cognitive, or mental limitations, needs assistance with heavy laundry such as carrying large loads of laundry.
0 0 2 - Unable to do any laundry due to physical limitation or needs continual supervision and assistance due to cognitive or mental limitation.
0 0 3 - Unable to do any laundry due to physical limitation or needs continual supervision and assistance due to cognitive or mental limitation.
0 0 4 - Unable to do any laundry due to physical limitation or needs continual supervision and assistance due to cognitive or mental limitation.
0 0 5 - Unable to do any laundry due to physical limitation or needs continual supervision and assistance due to cognitive or mental limitation.
0 0 6 - Unable to do any laundry due to physical limitation or needs continual supervision and assistance due to cognitive or mental limitation.
0 0 7 - Unknown

Key:
- More than one response allowed
- Only one response allowed

SOC Assessment Incorporating OASIS-BI (10/96)
## COMPREHENSIVE START OF CARE ASSESSMENT

(Also used for Resumption of Care Following Inpatient Stay)

### Medications

**Management of Oral Medications:** Patient's ability to prepare and take all prescribed oral medications, reliability and safety, including administration of the correct dosage at the appropriate time intervals. Excludes injectable and IV medications. **NOTE:** This refers to ability, reliability, compliance or diligence.

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
</tr>
</thead>
<tbody>
<tr>
<td>O O O</td>
<td>Able to independently take the correct medication(s) and proper dosage(s) at the correct times.</td>
</tr>
<tr>
<td>O O O</td>
<td>Able to take medication(s) at the correct times if: (a) individual dosages are prepared in advance by another person. OR (b) given daily reminders.</td>
</tr>
<tr>
<td>O O O</td>
<td>Unable to take medication unless administered by someone else.</td>
</tr>
<tr>
<td>O O O</td>
<td>Unable to take medications prescribed.</td>
</tr>
<tr>
<td>UK</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

**Management of Intravenous Medications:** Patient's ability to prepare and take all prescribed intravenous medications (infusions, enteral dose devices) reliably and safely, including administration of the correct dosage at the appropriate time intervals. Excludes all other forms of medication (oral tablets, injectable and IV medications).

<table>
<thead>
<tr>
<th>Prior</th>
<th>Current</th>
</tr>
</thead>
<tbody>
<tr>
<td>O O O</td>
<td>Able to independently take the correct medication and proper dosage at the correct times.</td>
</tr>
<tr>
<td>O O O</td>
<td>Able to take medication at the correct times if: (a) individual dosages are prepared in advance by another person. OR (b) given daily reminders.</td>
</tr>
<tr>
<td>O O O</td>
<td>Unable to take medication unless administered by someone else.</td>
</tr>
<tr>
<td>O O O</td>
<td>Unable to take medications prescribed.</td>
</tr>
<tr>
<td>UK</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

**Medication List:**

- O = more than one response allowed
- R = only one response allowed

**SOC Assessment incorporating OASIS-B1 (10/03)**
COMPREHENSIVE START OF CARE ASSESSMENT
(Also used for Resumption of Care Following Inpatient Stay)

(0010) Management of Injectable Medications: Patient's ability to prepare and take all prescribed injectable medications reliably and safely, including administration of correct dosage at the appropriate time/interval. Excludes IV medications.

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Able to independently take the correct medication and proper dosage at the correct times.</td>
</tr>
<tr>
<td>1</td>
<td>Able to take injectable medication at correct times if:</td>
</tr>
<tr>
<td></td>
<td>(a) Individual syringes are prepared in advance by another person.</td>
</tr>
<tr>
<td></td>
<td>(b) Given daily reminders.</td>
</tr>
<tr>
<td>2</td>
<td>Unable to take injectable medications unless administered by someone else.</td>
</tr>
<tr>
<td>NA</td>
<td>No injectable medications prescribed.</td>
</tr>
<tr>
<td>UK</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

(0015) Patient Management of Equipment: Excludes ONLY oxygen, Oral/nasal/nasopharyngeal nutrition equipment or supplies.

Patient's ability to set up, monitor and change equipment reliably and safely, add appropriate fluids or medication, characterize/describe of equipment or supplies using proper technique. (NOTE: This refers to ability, not compliance or willingness.)

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Patient manages all tasks related to equipment completely independently.</td>
</tr>
<tr>
<td>1</td>
<td>If someone else sets up equipment (i.e., portable oxygen tank, provides patient with prepared solutions), patient is able to manage all other aspects of equipment.</td>
</tr>
<tr>
<td>2</td>
<td>Patient requires considerable assistance from another person to manage equipment, but independently completes portions of the task.</td>
</tr>
<tr>
<td>3</td>
<td>Patient is only able to monitor equipment (e.g., liter flow, fluid in bag) and must call someone else to manage the equipment.</td>
</tr>
<tr>
<td>4</td>
<td>Patient is completely dependent on someone else to manage all equipment.</td>
</tr>
<tr>
<td>NA</td>
<td>No equipment of this type used in care</td>
</tr>
<tr>
<td>UK</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

(0020) Caregiver Management of Equipment: Excludes ONLY oxygen, Oral/nasal/nasopharyngeal nutrition equipment or supplies.

Caregiver's ability to set up, monitor, and change equipment reliably and safely, add appropriate fluids or medication, characterize/describe of equipment or supplies using proper technique. (NOTE: This refers to ability, not compliance or willingness.)

<table>
<thead>
<tr>
<th>Code</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Caregiver manages all tasks related to equipment completely independently.</td>
</tr>
<tr>
<td>1</td>
<td>If someone else sets up equipment, caregiver is able to manage all other aspects.</td>
</tr>
<tr>
<td>2</td>
<td>Caregiver requires considerable assistance from another person to manage equipment, but independently completes significant portions of the task.</td>
</tr>
<tr>
<td>3</td>
<td>Caregiver is only able to complete small portions of task (e.g., administer nebulizer treatment, characterize/describe of equipment or supplies)</td>
</tr>
<tr>
<td>4</td>
<td>Caregiver is completely dependent on someone else to manage all equipment.</td>
</tr>
<tr>
<td>NA</td>
<td>No caregiver</td>
</tr>
<tr>
<td>UK</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

ADDITIONAL ASSESSMENT INFORMATION DOCUMENTED ON OUTCOMES PLANNER VISIT #1

Signature of Nurse/Therapist: ____________________________ Date: _______________

Signature of Data Entry Clerk: ____________________________ Date: _______________
The CHAIRMAN. Vicki.

Ms. GIELLA. My name is Vicki Giella of the Area Agency on Aging in Northeastern Vermont. I am speaking from the perspective of a very small local rural agency, and I would just like to speak on behalf of balance in the area of individual need for privacy and agency needs to collaborate.

We work very much within an environment of confidentiality and releases. We never share information without release from clients; however, more and more as the health care environment changes we are working within coalitions and within consortiums of health care providers. So whatever—however the laws are changed, it is important that these local providers, and this is even within an area where there is not very much managed care, still in order to effectively work with patients, we are going to need to be able to share patient information. And so whatever kinds of ways in which the rules are changed and privacy is enhanced, we still—we do not want those rules to create a situation in which we will not be able to care effectively for very—in many cases very frail people and have their health care harmed by those regulations. I am saying this completely within the context of confidentiality and releases, personal releases.

The CHAIRMAN. Thank you, all four of you. I appreciate that information, and we will make that part of the record. We will be back in touch with all of you I am sure, as we proceed along. Thank you again.

Our final panel is Patrick Biggam of Vermont Labor Council; Brian Travis, M.D., medical director of Central Vermont PHO; Julie McGowan, UVM and AMIA.

Patrick first.

STATEMENTS OF PATRICK BIGGAM, ESQ., VERMONT LABOR COUNCIL, MONTPELIER, VT; BRIAN TRAVIS, M.D., MEDICAL DIRECTOR, CENTRAL VERMONT PHO, BERLIN, VT; AND JULIE MCGOWAN, UNIVERSITY OF VERMONT, AMIA, BURLINGTON, VT

Mr. BIGGAM. Thank you very much, Senator. My name is Patrick Biggam, and I am an attorney in Montpelier here on behalf of the Vermont Labor Council, AFL/CIO. I am a practicing attorney for 20 years in Montpelier, and our firm probably does more Worker's Compensation claims than any other firm in the State. In addition, I am past president of the Vermont Trial Lawyers Association.

The Vermont Labor Council's position is that protection of medical records is obviously a very good thing, and they are interested in the same rights and protections for the members as they are for the people and the public at large. The problem that we see happening, however, is Worker's Compensation carriers seek to be excluded from this bill. They have done so on the State levels, and they are doing so on the national level as well.

Now this exclusion is not a small exclusion. There are over 113 million employees covered by the Worker's Compensation system in this country. It is a $70 billion industry. In Vermont alone there are in excess of 25,000 first reports of injury. We represent granite workers obviously being in Central Vermont, but the bulk of our
clients are either secretaries, nurses aides, people in white collar positions as well as blue collar jobs. We have a lot of carpal tunnel injuries that we see. We have a lot of back injuries that we see that are related to all walks of life.

What happens when a person is injured in Vermont under the Worker's Comp system is they must sign a medical release for all of their medical records to the Worker's Compensation carrier. I have two releases that I have brought with me from two cases that we currently have. The first release is from Chubb Insurance Company, and it required our client to authorize every doctor, dentist, medical practitioner, pharmacist, hospital, insurance company, consumer reporting agency and employer to provide them with any information which is with respect to their diagnosis, treatment or prognosis of their medical condition and any nonmedical information about me which is relevant and requested. Now that is one type of release that is mandated by Chubb in order for a worker to collect Worker's Comp insurance.

The second release that a client was required to sign is even better. This one says, dear doctor, we have received notice of an industrial claim. Please send us all of your medical records. Since this is a Worker's Comp claim, no medical authorization is required. Now this was from Safeguard. What is happening in Worker's Comp?

The CHAIRMAN. Is that accurate?

Mr. BIGGAM. Well, in Vermont that is not accurate. The problem in Vermont is that there is a Form 7 release. Very few insurance companies provide the Form 7 to the worker. Of the 25,000 claims that are filed in Vermont, less than 1,500 have an attorney who gives any advice to the people about what their requirements are. And so as a result, they get away with a lot of these letters and releases.

The other thing is what are they doing with this information. We have talked about Big Brother and the collection of databases. I am informed that the American Insurance Services Group, which is an insurance company database, has in excess of 57 million claims in the national database, which I am sure include my clients and the records that I have been required to give to those insurance companies. They do it on the allegation of prevention of fraud. On the other hand, how many people want their medical records in a national database of this sort?

So basically I would argue and simply in a presentation is we support the bill. We would not like to see any exclusion for Worker's Compensation because what you are trying to protect on one hand is going to be let out the back door on the other if Worker's Comp is excluded. Thank you.

The CHAIRMAN. Thank you.

Dr. TRAVIS. My name is Brian Travis. I am the medical director at the Central Vermont PHO. I am also the director of the department of laboratory, medicine and pathology both at Central Vermont Hospital and Gifford Memorial Hospital in Randolph. As a specialist in surgical pathology, I serve around 25,000 people in the Central Vermont area and render 40 or 50 diagnoses—individual diagnoses a day at a subspecialty level on these patients which influence their life profoundly including whether or not they are
going to have major surgery, whether they have to undergo dangerous chemotherapy or radiation therapy.

I am sitting there like in my office, which is right upstairs, every day making these huge decisions, and I need to have information right at my hands to important X-rays, to previous diagnoses of cancer or not, to the fact of whether they might have HIV or any other thing that might help me to make this decision that influences their life profoundly. A mistake made there can lead to absolute tragic consequences.

And subspecialists like myself need to have as much information as we need. I do not care what their name is, but we need to have as much information as we need at the point that we make that decision if you want us to continue to support the infrastructure of the best medical system in the world.

Along with this kind of access needs to go—needs to be accompanied by protection which is the whole thrust of your legislation. The audit system alluded to by Dr. Gettinger and later by Dr. Ward is something that every hospital that I know of in Vermont is capable of implementing if they have the right mandate. Every time a medical record is accessed by a provider, there can be a log of who it was and when they did it. And if we were mandated, we could go through these logs in a systematic quality manner to look at the accesses that were not appropriate and take appropriate action.

A guideline for appropriate action also needs to be promulgated. It is left up to the individual institution what is appropriate action at this point. The reason I know this is because I am a computer programmer also in the modern computer languages and am familiar with the systems that we have here, and at Gifford and in other places.

The last vignette that I have for you is that it is a Federal mandate through the JCAH, or joint commission, to put the patient’s name on all requisitions for lab tests or anything else. That needs to be done away with so that maybe we could use a number or something so that six or eight people do not see the name as this request is entered into the system.

The CHAIRMAN. Thank you. Very helpful.

Yes. Julie, please proceed.

Ms. McGOWAN. My name is Julie McGowan, and I am a medical informatics researcher at the University of Vermont. I am also a member of the Public Policy Committee of the Medical Informatics Association, a national association of almost 4,000 physicians, researchers, health information professionals and others interested in the use of information technology to support health care delivery. My comments reflect my own and as a member of this organization.

I would like to begin by thanking Senator Jeffords for introducing S. 578 and both senators for taking a leadership role in examining one of the most critical issues facing health care today. I would also like to thank the staff of the U.S. Senate committee for their willingness to listen to our concerns and formulate a balanced approach to the need for patient confidentiality and the need for information dissemination in support of health care delivery and research.
Information professionals are commonly perceived as placing the need for easy transmission of health care information ahead of the privacy rights of patients. Nothing could be farther from the truth. We support strong uniform Federal confidentiality standards to ensure that each individual can feel secure that his or her health information is protected. We also feel that all health information should receive the same protections regardless of format, whether that format is a paper record or an electronic medical record.

In the area of research, academic health centers such as the one at the University of Vermont have established thorough IRBs to review proposed research and ensure that patient’s rights are protected. This is especially critical in prospective research in which patients are recruited to participate in various studies. However, sometimes it is more effective to use data from large aggregate databases or data elements from individual records from which all identifying information has been removed.

In these instances it is—it frequently becomes problematic to seek individual patient permission to use information in a medical record which is 10 or 20 years old. Although the information is old and/or aggregated, it is still critical to ensure the patient’s privacy. Again, the IRB should review such study designs with the concept of patient confidentiality protection as paramount.

In today’s health care environment high quality and low cost are top priorities. Institutional quality assurance initiatives also need access to patient records and aggregate databases. Often these do not fall under the umbrella of research or IRBs. However, it is essential that all policies and procedures regarding the use of protected health information be consistent with Federal policy for the protection of human subjects.

Senator Jeffords’ bill has gone a long way in addressing these issues. It presents a balanced approach between the rights of patients regarding personal health information and the needs of health care providers and researchers to use such information to deliver care to the individual and to promote health for all citizens.

Thank you, Senator Jeffords.

The CHAIRMAN. Well, thank you all very much. The time has come to an end, unfortunately. In many respects, we have heard wonderful testimony today. I deeply appreciate everyone who has assisted us. I also, of course, want to thank the Central Vermont Medical Center, especially its CEO, Daria Mason, who gave us fantastic help, I appreciate it; my reporter who is still rapping away and doing very well; and all of you for coming and helping us with this very, very important issue. The outpouring of people we have had today, as well as the testimony from our witnesses, demonstrates that this is obviously a matter of deep and high concern to Vermonters.

I am sure I speak for Senator Leahy when I say that he and I will make sure that Vermont’s interests are as well protected as possible. Thank you all. With that we will close the hearing.

The record will be open for 2 weeks if anyone desires to correspond with us.

[Whereupon, at 12:16 p.m., the committee was adjourned.]