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# Keeping our patients' secrets

## ■ ABSTRACT

Protecting the privacy of the patient's medical record is a central issue in current discussions about a patient bill of rights, and controversy over a proposed "unique health identifier" has raised the decibel level of these discussions. At the heart of the debate is how best to resolve the inherent conflict between the individual's right to privacy and the need for access to patients' health information for reasons of public health, research, and health care management.

**D**EBATE ABOUT A "UNIQUE HEALTH IDENTIFIER" has refocused public attention on the privacy, confidentiality, and security of medical records.<sup>1</sup>

The unique health identifier is part of a national health care identification system required by the Health Insurance Portability and Accountability Act of 1996 (HIPAA). Many believe it threatens the privacy, confidentiality, and security of medical records; others see it as necessary to optimal public health, epidemiologic research, and management of care.

Unfortunately, in the real world, patient privacy and confidentiality are broken regularly, even without any increased vulnerability to breach of confidence that a mandated unique health identifier might impose.<sup>2,3</sup> These breaches benefit no one, and physicians cannot condone them.<sup>4</sup> Our time-honored, loosely controlled medical record system frequently fails to protect the individual's right to privacy, ignores confidentiality, and offers little security.<sup>5,6</sup> In addition, the openness of the academic environment may conflict with patient confidentiality. Electronic data technology may facilitate these violations of confidentiality, but it is not the real villain.<sup>7</sup>

## ■ HOW DO PRIVACY, CONFIDENTIALITY, AND SECURITY DIFFER?

The definitions of privacy, confidentiality, and security used here are those commonly used in the professions concerned with these issues:

**Privacy** is the limitation of awareness of personal information to the patient and possibly a few others whom the patient selects. Privacy is governed by the patient alone. Private information may or may not be recorded in the medical record; if it is, then it is regarded as confidential.

**Confidentiality** is the limitation of awareness of private information to those who must have it in order to provide desired service to the patient. Confidentiality is governed by the "need-to-know" principle. Through disclosure and formal authorization to release information, the decision about who needs to know private information is controlled to some extent by the patient. Information in the medical record is presumed to be confidential.

Private does not always mean confidential, however. Although similar, the concepts of privacy and confidentiality are not necessarily congruent. Information considered private by the patient should usually be held confidential by those to whom it is entrusted, but not always. For example, a person with syphilis may regard that information as *private*, but by law it must be reported; therefore, it is not *confidential*.

**Security** refers to safeguards to the confidentiality and integrity of recorded information. In health care, such protections have traditionally been loose, ostensibly to promote ease of information sharing among those who need to know it.

**A person with syphilis may regard that fact as private, but by law it must be reported, so it is not confidential**



## ■ MANAGED CARE NEEDS ACCESS TO CONFIDENTIAL INFORMATION

In the rush to control health care costs, attention has focused on managing delivery of care by using increasingly controversial tools collectively referred to as managed care.<sup>8</sup> Patients have become the consumers, doctors and hospitals are now the providers, and third-party payers, including the government, are the customers.

In this uneasy ménage à trois, providers find themselves caught between the often competing interests of patients and payers. This situation raises significant ethical issues. Patients want their privacy respected. Providers are, for the most part, responsible for keeping such information confidential, thus protecting patient privacy. Yet payers and other interested parties such as researchers and public health workers<sup>9</sup> may want the information. Their purposes might include, for example, monitoring immunization, disease screening rates, or compliance with treatment protocols. Because of the increasing use of computers to store medical data, such information is much easier to assemble and disseminate than in the past. As access to patient data becomes technically easier, patients are becoming progressively uneasy.

Passage of the HIPAA intensified physician concern because of its severe legal penalties for unauthorized release of medical data. The potential passage of a patient bill of rights promises more of the same.

## ■ THE PUBLIC NEEDS ACCESS TO CONFIDENTIAL INFORMATION

The need to protect the public against threats to health or safety conflicts with the need for confidentiality of health information.<sup>5,10</sup> It is a conflict between the needs of the many<sup>11,12</sup> and the rights of the individual<sup>13</sup>—an ethical issue that has concerned physicians since the time of Hippocrates.

The issue is particularly poignant in the case of deadly infectious diseases such as AIDS,<sup>14</sup> mental health issues potentially affecting public safety,<sup>15</sup> criminal investigations,<sup>16</sup> and certain disastrous genetic conditions whose transmission to the next generation could be prevented.<sup>17</sup>

## ■ OTHERS NEED ACCESS TO CONFIDENTIAL INFORMATION

Other examples where an individual or entity has or perceives a compelling need to obtain confidential medical information include:

- The “right” of family members to know genealogic information of interest, such as biological parentage<sup>18</sup>
- The “need” of an employer to know an injured employee’s readiness to work<sup>19</sup>
- The “need” of a victim to have the assailant tested for transmissible disease
- The desire of insurance companies to know medical information for coverage and underwriting purposes.<sup>20</sup>

History suggests that the list of such examples will expand as medical understanding produces more uses for the information that can be found in personal medical records.

## ■ LEGISLATION AND THE UNIQUE HEALTH IDENTIFIER

### Health Insurance Portability and Accountability Act of 1996 (HIPAA, PL 104-101)

The HIPAA mandates six requirements for patient data. The requirements include:

- Ability to transmit health information electronically
- Unique health identifiers for individuals, employers, health plans, and providers
- Code sets and classification systems for each data element in electronic health care transactions
- Security of health care information systems
- Procedures for electronic transmission and validation of signatures
- Data elements needed to coordinate benefits and process claims.<sup>21</sup>

In August 1998, the Health Care Financing Administration proposed rules for implementing these requirements, with a comment period that ended in October 1998. Interestingly, the deadlines for finalizing some of these rules had already passed when the draft rules were published. Final rules have not yet been promulgated.

### Options for the unique health identifier

The unique health identifier was viewed as

**Managed care puts providers in the middle of patients’ and payers’ competing interests**

essential to administrative simplification, and was expected to result in better quality of care, lower administrative costs, and other benefits.

Options for the unique health identifier were subjected for public comment in July 1998.<sup>1</sup> It was an eye-opener. Those options included a numerical identifier based on the Social Security number, a totally new numerical identifier not based on the Social Security number, a biometric identifier based on physical attributes (eg, fingerprints, retinal pattern analysis, iris scan, voice pattern, DNA analysis), and a numerical identifier based on the Civil Registration System (eg, birth files, visas, “green cards”). Additional proposals recommended methods that would allow cross-linking of medical records across systems without a unique personal identifier.

Some of the more exotic options such as iris scan or DNA analysis smacked of a Big Brother and other predictions of paranoid futuristic fiction.

The unique health identifier would allow individuals to be tracked throughout their use of the health care system.<sup>22</sup> As in some other countries, it could permit the establishment of a state or national health database,<sup>23–25</sup> which would be useful not only for keeping track of utilization, but also for public health and research.<sup>26</sup>

#### **Government’s good intentions greeted with skepticism**

To say opinion on the unique health identifier is deeply divided<sup>1</sup> may be the understatement of the year. The tracking ability that such an identifier would give the government—or any organization with access to it—has Big Brother overtones. General confidence in the good intentions of government agencies interested in controlling costs would go a long way toward defusing public opposition, but such confidence does not appear to exist now. The public outcry resulted in suspension of the attempt to define a unique health identifier.

#### **Research and the unique health identifier**

People are also concerned about the use of identifiable data in research pertaining to epidemiologic,<sup>27</sup> genetic,<sup>28</sup> and public health,<sup>29</sup> and requirements for consent have been revisited and strengthened.<sup>12,30,31</sup> People have even

raised doubts about the use of statistically analyzed, aggregated data stripped of personal identifiers.<sup>32,33</sup> There is little trust in the integrity of the whole clinical research apparatus, and the patient bill of rights and the HIPAA address these misgivings in proposals for providing guarantees about informed consent for the release of data. Other countries have been more sanguine about trading off some confidentiality for the perceived public good.<sup>34</sup>

#### **■ SECURITY ISSUES**

Although a detailed discussion of the fine points of data security is beyond the scope of this paper, the topic is important ethically,<sup>35</sup> and medical record security is necessary to meet regulatory requirements.<sup>36</sup>

Security protects both the confidentiality and the integrity of data.<sup>37</sup> The growth of telemedicine complicates the issue of security.<sup>38</sup> Military-type security measures and standard database protection methods are inappropriate, however, because they prevent ready access to data, which is essential for efficient delivery of medical care.<sup>39</sup>

Currently available security methods include:

- User validation and access control, including secure electronic signatures<sup>40</sup>
- “Depersonalization” of data with a secure identifier control facility acting as a network file access table
- A “virtual record” instead of an integrated file, which would be reconstructed at the time of access<sup>41</sup>
- Encryption of data.<sup>42,43</sup>

In Europe, standards for health data security have been promulgated.<sup>44</sup> Grotan and Iverson<sup>45</sup> pointed out the inadequacies of present day safeguards on medical data. This was reemphasized in a 1997 report of the National Research Council of the National Academy of Sciences.<sup>40</sup> In the United States, no system has yet been widely recognized as ideal, although several are under investigation.<sup>46,47</sup>

Even though strong security procedures are in place to protect confidential medical information, individuals are regularly asked to waive confidentiality (eg, when applying for insurance), and few controls on dissemination exist once the waiver is signed.

**Handle confidential information as you would want your own medical information handled**



## ■ WHAT IS THE MESSAGE FOR PHYSICIANS?

Careful attention to confidentiality and security of patient-related information, especially the medical record (whether paper or electronic), is an absolute must.

An adequate review mechanism is needed, possibly involving the physician, to ascertain that requests are legitimate before information is released.

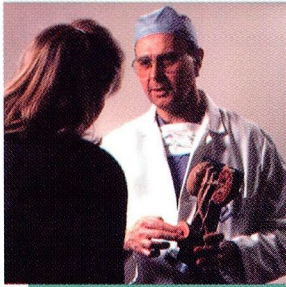
Totally aside from any threat to confidentiality posed by converting medical records to electronic form or using a unique health identifier, discussing sensitive information in public places such as elevators and cafeterias is all

too common and must stop.<sup>2</sup> A simple standard is to handle confidential information the way you would want your own medical information handled. Accrediting agencies, such as the Joint Commission on Accreditation of Health Care Organizations,<sup>48</sup> have developed formal standards that should help office-based physicians devise specific, useful approaches.

Physicians and other health care providers will increasingly be held accountable for breaches of confidentiality, and the penalties, not to mention the loss of patients' trust, will be severe. Beyond that, we owe it to our patients to respect and protect their confidences. Only then can the trust that underlies the doctor-patient relationship survive. ■

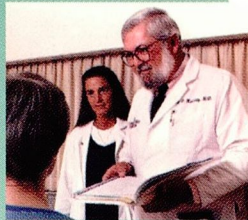
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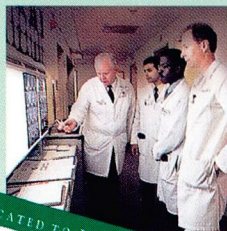
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