1.2.9 Use of Remote Sensing and Monitoring Devices

Sensing and monitoring devices can benefit patients by allowing physicians and other health care professionals to obtain timely information about the patient’s vital signs or health status without requiring an in-person, face-to-face encounter. Implantable devices can also enable physicians to identify patients rapidly and expedite access to patients’ medical records. Devices that transmit patient information wirelessly to remote receiving stations can offer convenience for both patients and physicians, enhance the efficiency and quality of care, and promote increased access to care, but also raise concerns about safety and the confidentiality of patient information.

Individually, physicians who employ remote sensing and monitoring devices in providing patient care should:

(a) Determine whether using one or more such devices is appropriate in light of individual patients’ medical needs and circumstances, including patients’ ability to use the chosen device appropriately.

(b) Explain how the device(s) will be used in the patient’s care and what will be expected of the patient in using the technology, and disclose any limitations, risks, or medical uncertainties associated with the device(s) and data transmission.

(c) Obtain the patient’s or surrogate’s informed consent before implementing the device in treatment.

Collectively, physicians should:

(d) Support research into the safety, efficacy, and possible non-medical uses of remote sensing and monitoring devices, including devices intended to transmit biometric data and implantable radio frequency ID devices.

(e) Advocate for appropriate oversight of remote sensing and monitoring devices.

AMA Principles of Medical Ethics: I,III,V

Background report(s):

CEJA Report 3-A-16 Modernized Code of Medical Ethics
CEJA Report 5-A-07 Radio frequency ID devices in humans
1.2.9 Use of Remote Sensing and Monitoring Devices

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Individually, physicians who employ remote sensing and monitoring devices in providing patient care should:

(a) Determine whether using one or more such devices is appropriate in light of individual patients’ medical needs and circumstances, including patients’ ability to use the chosen device appropriately. [new content addresses gap in current guidance]

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AMA Principles of Medical Ethics: I,III,V
INTRODUCTION

Radio frequency identification (RFID) tags are computer chips connected to miniature antennae that can be used to transmit information electronically via a proximate RFID reader. The use of these devices in health care represents another promising development in information technology, but also raises important ethical, legal and social issues. Specifically, the use of RFID labeling in humans for medical purposes may improve patient safety, but also may pose some physical risks, compromise patient privacy, or present other social hazards.

This report responds to Resolution 6 (A-06), “RFID Labeling in Humans,” which called for study of the medical and ethical implications of RFID chips in humans. This report focuses on ethical issues in the use of RFID chips, specifically in regard to their implantation for clinical purposes.

BACKGROUND

Radio frequency identification devices utilize wireless technology to communicate data via signals in the radio frequency range of the electromagnetic spectrum. Data are stored in a microchip attached to an antenna, and packaged so that they can be attached to or embedded in products, animals, or people.

The two main types of RFID tags are passive and active. Passive tags contain no internal power supply. They convert the radio frequency energy emitted from a reader device into signals that transmit stored data for a distance of a few feet. These passive devices currently have restricted amounts of data storage and are of limited functionality, because the information they contain cannot be modified.

In comparison, active RFID tags contain an internal battery, which provides increased reliability, longer transmission ranges, on-tag data processing and greater data storage. While their capacity...
to process data internally allows for expanded capabilities in the future, their greater transmission
range presents a more substantial threat to data confidentiality and patients’ privacy.
In October 2004, the US Food and Drug Administration (FDA) approved the first RFID tags
specifically intended for human implantation.\(^2\) Approved RFID devices are currently limited to
passive units, intended for identifying patients. Active RFID chips may be approved in the future.

Human-implanted passive RFID devices that identify patients may also contain essential biometric
and medical information. The tags are primarily intended for patients with chronic diseases, such
as coronary artery disease, chronic obstructive pulmonary disease, diabetes mellitus, stroke or
seizure disorder, or are implanted into patients with medical devices such as pacemakers, stents, or
joint replacements. These devices are approximately the size of a grain of rice, and are implanted
under the skin via a hypodermic-type needle in less than one minute.\(^3\)

RFID tags may promote the timely identification of patients and expedite access to their medical
information. As a result, these devices can improve the continuity and coordination of care with
resulting reduction in adverse drug events and other medical errors.\(^4\)

RFID tags also may improve efficiency within the health care system. In conjunction with
improved medical record management, these devices may facilitate access to patient records,
medication lists, and diagnostic tests.\(^5\) To be maximally effective, however, the information in
these devices must be adequately integrated into present clinical information and communications
systems, laboratory databases, and pharmacy systems.\(^1\)

Appropriate processes also must be developed to inscribe, read and archive data stored on RFID
tags. As new designs enter the marketplace, the emergence of competing standards may present
problems for hospital staff if a patient’s ID tag proves incompatible with the interrogation devices
employed by the hospital.\(^1\)

These devices may present physical risks to the patient. Though they are removable, their small
size allows them to migrate under the skin, making them potentially difficult to extract. However,
this tendency may be minimized by constructing RFID tags from materials that permit surrounding
tissue to encase the device. In addition, RFID tags may cause electromagnetic interference, which
may interfere with electrosurgical devices and defibrillators.\(^1\) Finally, it has not been determined
whether RFID tags might affect the efficacy of pharmaceuticals.\(^1,6\)

The primary concerns surrounding human RFID labeling pertain to their potential impact on patient
privacy and security. Physicians must assure patients that their medical information will be held in

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confidence (see Opinion E-5.05, “Confidentiality”). Moreover, maintenance of privacy is required
to protect patients from embarrassment, potential social discrimination, loss of health care
coverage, or other detrimental consequences (see Opinion E-5.059, “Privacy in the Context of
Health Care”).

At this time, the security of RFID devices has not been fully established. Physicians, therefore,
cannot assure patients that the personal information contained on RFID tags will be appropriately
protected. In light of these security concerns, the FDA currently requires RFID transponders to
store only a unique electronic identification code to be read by the scanner. This identification
code can then be used to access patient identity and corresponding health information stored in a
database.

To protect confidentiality and privacy, the medical community should advocate for the adoption of
other protections, such as computer encryption or digital signatures. Ultimately, the medical
community should undertake appropriate efforts to prevent unauthorized access to patients’
information contained on RFID tags (see also E-5.07, “Confidentiality: Computers,” AMA Policy
Database).

INFORMED CONSENT

To properly respect patient autonomy, RFID tags should not be implanted or removed without the
prior consent of patients or their surrogates (see E-8.08, “Informed Consent,” and E-8.081,
“Surrogate Decision Making”). During the consent process, decision-makers should be informed
of the potential risks and benefits associated with RFID tags, including the many uncertainties
regarding their efficacy. Patients are also entitled to know who will be granted access to the data
contained on RFID tags and the purposes for which this information will be used.

FURTHER CONSIDERATIONS

It seems likely that utilization of RFID devices for medical purposes will expand. The medical
profession must continue to monitor the efficacy of these devices. If RFID tags are proven to
benefit patient care significantly, the profession should advocate for widespread adoption of RFID
technology, and for policies that make RFID tags available to all patients who would benefit (see

However, if objective evidence demonstrates negative consequences that outweigh the benefits in
relation to health care, the medical profession will bear an important responsibility to oppose the
use of RFID labeling in humans.

Finally, physicians should be aware of emerging non-medical applications of human-implantable
RFID devices. For instance, active RFID technologies might be considered for the tracking or
surveillance of individuals who pose a threat to others. Although this is only one of many possible
uses of RFID technology in the future, it alerts the medical profession to the need for continuous
assessment of the appropriate role of physicians participating in RFID labeling of human beings. Indeed, certain uses could constitute an infringement upon patients’ individual liberties, placing physicians in a position to act as patient advocates by promoting the use of other, less intrusive alternatives, when available.

CONCLUSION

RFID technology has the potential to improve patient care as well as patient safety. However, the safety and efficacy of human-implantable RFID devices has yet to be established. Therefore, the medical community should support further investigations to obtain the data necessary to make informed medical decisions regarding the use of these devices. The medical community should also be sensitive to potential social consequences of RFID devices, such as non-medical applications in law enforcement.

RECOMMENDATION

The Council on Ethical and Judicial Affairs recommends that the following be adopted and the remainder of the report be filed.

Radio frequency identification (RFID) devices may help to identify patients, thereby improving the safety and efficiency of patient care, and may be used to enable secure access to patient clinical information. However, their efficacy and security have not been established. Therefore, physicians implanting such devices should take certain precautions:

1. The informed consent process must include disclosure of medical uncertainties associated with these devices.
2. Physicians should strive to protect patients’ privacy by storing confidential information only on RFID devices with informational security similar to that required of medical records.
3. Physicians should support research into the safety, efficacy, and potential non-medical uses of RFID devices in human beings.

(NEW HOD/CEJA Policy)

Fiscal Note: Staff cost estimated at less than $500 to implement.
REFERENCES

1 Ingeholm; Mun, K; Mun, SK. RFID in Healthcare: The Applications, and Obstacles, Are Many; *Journal of AHIMA*; 2006. 77(8): 56-62.


3 DeNoon D. Chip implants: Better care or privacy scare. 2005. Accessible at”
   http://www.webmd.com/content/Article/109/109216.htm

4 Wicks, AM; Visich, JK; Li, Suhong. Radio Frequency Identification Applications in Hospital Environments; *Hospital Topics*.2006; 84(3): 3-8.

5 VeriMed™ Information Center for Patients; http://www.verimedinfo.com/content/intro/physicians

6 Wasserman, Elizabeth. A Prescription for Pharmaceuticals; *RFID Journal*. 2006. Accessible at:
   http://www.rfidjournal.com/magazine/article/1739/1/173/