Non-clinical Use of Obstetric Ultrasound: Medico-Legal Implications in the USA

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The United States is known as the land of free enterprise, so it should be no surprise that medical equipment has undergone a transformation to use by laypersons for commercial reasons. Eighty years ago the US Patent Office granted a patent to Dr. Jacob Lowe, a Boston physician, for a fluoroscopic device to aid in fitting shoes. The doses of radiation to which untold numbers of children and adults were exposed is not precisely known, but may have been as high as 7-14 Rads for a 20 second exposure\(^1\).

Shoe-fitting fluoroscopes were regulated out of existence by the 1950s. There are a host of other non-medical uses of sophisticated technology available on the patient’s initiative, however. CT of the heart is recommended for anyone fitting a variety of “risk factors” by one center. These risk factors include men over 35 and women over 40, high stress levels and a sedentary lifestyle\(^2\). One company provides pricing on line, with a coronary artery scan for $395 in Los Angeles or Chicago. For some reason the same scan costs $100 more in Washington, DC\(^3\). Full body scans are also available. Insurance, the consumer is warned on these sites, does not cover these scans, although credit cards, checks or cash are welcome.

Coronary artery and full body CT scans use expensive equipment, and potentially dangerous ionizing radiation, unlike ultrasound. So, the development of a booming business of prenatal ultrasound entertainment centers in the US might have been predictable. With names like “Womb With a View”, “Fetal Fotos”, and “Baby Insight”, these offices provide 3D and 4D sonography on demand, with convenient hours and packages including your baby’s movements set to music on a CD, and comfortable theater style seating for guests. Perhaps worst of all, some offer reports of “limited medical ultrasounds”.

Other companies offer franchising information on the web, including business plans, legal forms, practice standards, and even physician oversight\(^4\).

At present there is no way of estimating the number of such businesses in the US. There are no standards for the scans that are
performed, no federal or state oversight of the offices offering these services, and no way of knowing who is performing the scans on what machines.

The US Food and Drug Administration classifies ultrasound machines as “Class II devices”, (21 CFR 892.1550, 1560 & 1570) subject to less stringent oversight than Class III devices, defined as those that support or sustain human life, are of substantial importance in preventing impairment of human health, or which present a potential, unreasonable risk of illness or injury. So, while ultrasound equipment comes with a label specifying that it is to be used by or on the order of a physician, the degree of oversight that the FDA expects to apply to the actual use of the machine is much less than for a device that produces ionizing radiation.

In fact, ultrasound machines, including 3D and 4D units, are readily available for purchase by anyone with sufficient cash on www.ebay.com. A recent search on that site (9/26/07) produced machines for purchase for as little as $15,000, though the listings for each available unit included a warning that credentials of the purchaser would be verified to ensure compliance with FDA regulations.

In 2005 the AIUM developed a statement on Keepsake Fetal Imaging\(^5\), written by a task force that I chaired and approved by the Board of Governors. The AIUM statement addresses the issue of qualifications of the individual performing the sonogram. The position of the AIUM is that only qualified individuals should perform sonograms, and that anyone performing fetal sonograms must be able to recognize important conditions such as birth defects, and conversely artifacts that may mimic fetal pathology. The lack of regulation of entertainment sonography, and the absence of standards for the performance of the sonograms makes it impossible to know what is being done.

While there are anecdotal reports of missed diagnoses during entertainment scans\(^6\), we have no systematic collection of data to understand the likelihood of missed diagnoses, or false positive diagnoses of anomalies. Similarly, we have no data to appreciate
whether significant numbers of previously unsuspected anomalies are being detected during entertainment ultrasounds in medical or non-medical facilities.

What we do know in this area is that there is no standard method of training the franchisees who open freestanding facilities. One franchiser includes over 20 bullet points on its website elucidating the support offered to franchisees, but none of the points include training to perform ultrasounds. Another offers protected territory and marketing assistance, as well as “comprehensive training in all areas of the business, including: Day-to-Day Operations; Employees; Bookkeeping and accounting procedures; Advertising and Marketing programs”, but not specifically technical assistance in how to scan fetuses. Unfortunately, while credentialed sonographers have opened some franchises, entrepreneurial businesspersons who do not share the professional training and attitudes expected of registered sonographers have opened others. One of these was quoted in the Wall Street Journal saying "I don't care if the fetus has three legs, I'd only point out two. I don't care if their uterus has fibroids, or if they have too much or too little amniotic fluid or where the placenta is. I have informed these people I'm not a doctor, that I'm not trying to find abnormalities.".

The issue of reports generated by freestanding facilities has not been heavily scrutinized. Sonographers, and especially non-sonographers who perform entertainment scans, are not licensed in any state to practice medicine independently, and the issuance of any report of an imaging study is usually considered to be the purview of licensed physicians. Thus far, states have not extensively pursued this, nor are there cases of civil suits related to missed diagnoses or bad outcomes after entertainment ultrasounds. One case in Texas attempted to use the complex and technical concept of changing the use of the ultrasound machine from the original labeling, thus making it a Class III device. The Appellate Court denied this part of the complaint, though the court upheld a misbranding claim for use of a prescription device without the order of a physician.

In 2005, also in Texas, the Attorney General reached an agreement
with four fetal imaging companies requiring physician oversight of entertainment ultrasounds. How this is being accomplished is not specified in the legal documents\textsuperscript{10}.

In California, in the wake of publicity regarding actor Tom Cruise’s revelation that he had purchased an ultrasound machine to observe his fiancée Katie Holmes’ fetus, the legislature considered and passed a bill in 2006 (AB 2360) making the sale of ultrasound machines to non-medical sites illegal. Although this might be considered something already prevented through FDA regulations, the legislature felt compelled to act. Governor Schwarzenegger vetoed the bill, saying that it conflicted with prior legislation requiring entertainment ultrasound sites to give clients written disclosure of the position opposing entertainment scans (AB 2049, 2004).

The AIUM Task Force on entertainment ultrasound examined professional codes of conduct to see if guidance in this area could be developed from existing statements. In preparing the statement, the committee considered the various settings in which entertainment scans might be performed. These included:

- In a physician’s office as part of a medically indicated scan
- In a physician’s office as a separate event and paid for by the patient outside of insurance payments for medical care
- In a freestanding commercial facility

The AIUM looked at standards from the American Medical Association and the American College of Obstetricians and Gynecologists. The AMA specifically comments on two types of products that might be offered in physicians’ offices for additional costs, health-relate and non-health-related. The policy provides that “physicians may sell low-cost non-health-related goods from their offices for the benefit of community organizations, provided that (1) the goods in question are low-cost; (2) the physician takes no share in profit from their sale; (3) such sales are not a regular part of the physician's business; (4) sales are conducted in a dignified manner; and (5) sales are conducted in such a way as to assure that patients are not pressured into making purchases.”\textsuperscript{11}
The stipulations on health-related goods are more restrictive. They include (excerpted from \cite{12}, deleted text indicated by […]):

“(1) Physicians should not sell any health-related products whose claims of benefit lack scientific validity. [Based…] on peer-reviewed literature and other unbiased scientific sources that review evidence in a sound, systematic, and reliable fashion.

(2) Because of the risk of patient exploitation and the potential to demean the profession of medicine, physicians who choose to sell health-related products from their offices must take steps to minimize their financial conflicts of interest. The following guidelines apply: (a) In general, physicians should limit sales to products that serve the immediate and pressing needs of their patients. For example, if traveling to the closest pharmacy would in some way jeopardize the welfare of the patient (eg, forcing a patient with a broken leg to travel to a local pharmacy for crutches), then it may be appropriate to provide the product from the physician’s office. […]

b) Physicians may distribute other health-related products to their patients free of charge or at cost, in order to make useful products readily available to their patients. […]

(3) Physicians must disclose fully the nature of their financial arrangement with a manufacturer or supplier to sell health-related products. […]

(4) Physicians should not participate in exclusive distributorships of health-related products which are available only through physicians' offices. […]”

The American College of Obstetricians & Gynecologists has adopted similar language to that of the AMA, saying “It is ethical and appropriate, however, to sell products to patients as follows: sale of devices or drugs that require professional administration in the office setting; sale of therapeutic agents, when no other facilities can provide them at reasonable convenience and at reasonable cost; sale of products that clearly are external to the patient–physician relationship, when such a sale would be considered appropriate in an external relationship; and sale of low-cost products for the benefit of community organizations.” \cite{13}
As the AIUM Task Force considered the options for scanning in physician offices, there were two possible ways to define the scan, as medical or non-medical procedures. If we define entertainment scans as non-medical, offering that service violates the guidelines of the AMA as they are not low-cost, the physician would gain profit from the transaction, and there is likely to be some subtle or overt pressure on the patient to have the scan. Similarly the ACOG guideline would be violated, as the scan would not be a low-cost item sold for the benefit of a community organization.

If the entertainment scans are defined as medical services, the AMA guidelines are violated because they do not serve immediate and pressing needs of patients, and are not offered free of charge or at cost. The ACOG guidelines would similarly seem to prohibit performing entertainment scans if they are defined as non-medical.

Conclusions

Where does this leave us? In the USA, the Food and Drug Administration has regulatory power over the manufacture and sale of ultrasound systems, but seems to have delegated the enforcement of violations of their rules to the states. The states are taking little or no action.

The market for entertainment scans has been driven in part by reluctance of some imagers to provide still or video images to patients. This has been due, at least in part, by concerns that images provided to patients could be used as evidence in claims of failure to diagnose a congenital abnormality. This argument has inherent weaknesses. Keeping a copy of any still images provided to the patient in the medical record covers at least part of this concern. Parallel video clips can also be retained. For some anomalies that develop later in gestation, for example duodenal atresia, the image provided to the patient may well help exonerate the physician.

Our practice is to give multiple still images to all patients having medically indicated scans, including 3D images if time permits and the fetus is in a good position. We also provide video clips to patients who bring their own tapes or discs. We do this in the belief that good
relations with patients are important, and a potential deterrent to future professional liability claims.

The American Institute of Ultrasound in Medicine is firm in its opposition to non-medical use of obstetric sonography, and the commercialization of fetal sonograms by non-professionals. We continue to encourage the FDA to enforce its regulations in this area. AIUM also encourages ultrasound equipment manufacturers to agree not to sell imaging systems to non-medical imaging facilities.
References

1. Oak Ridge Associated Universities web site

2. Princeton Longevity center web site

3. Heart Check America web site

4. United Imaging Partners web site


7. Womb with a view web site


10. State of Texas Attorney general web site
