Medical Device Safety & Innovation

preparing for system integration at the sharp edge of healthcare

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Medical Device Safety & Innovation

“Integrating the clinical environment” is essential to create error-resistant systems to improve patient care and healthcare efficiency
Current state

... at the sharp edge of high-acuity patient care ...
Clinical environments are crowded with complex, life-saving technology.
Forward-area Operating Room in Iraq

Critical Care Air Transport
Intensive Care Unit

Credit: T. Rausch
How do we connect medical devices to the EMR?

Hint: it isn’t “plug and play” like USB memory sticks
How will we interpret this electronic medical record?
Mass General Hospital
Operating Room of the Future (OR#49)
Lessons from the OR of the Future project

- Comprehensive integration of clinical and non-clinical devices and systems could provide:
  - **Workflow Support** require “closing the workflow loop”
  - **Smart Alarms** require “contextual awareness”
  - **Safety Interlocks** require tight system integration

Not limited to the OR: ICU, ER, home, etc.
Example of error resistant system:
Landing gear not down? -> **Smart ALARM**

Contextual awareness requires data from several device and systems.
“Hudson River Over-ride” - Augment awareness, not control
Examples of 2 clinical procedures and associated safety issues ->

(From our “Clinical Scenarios” Repository)
Scenario: Surgical Fires

600 surgical fires each year

The most severe burns are internal – in the lungs
Caused by burning breathing tubes
Airway Laser Surgery + O₂ -> Fire

• O₂ in respiratory gas supports combustion.
• If laser hits breathing tube, could produce devastating burn.
• Surgical team must “remember” to minimize O₂
A Solution: Laser-O₂ Interlock

- Monitor breathing O₂ concentration. (This is already measured.)
- Safety interlock to prevent laser activation if O₂ > 25%

Proposed and published in 1999!

NOT Commercially available
Scenario:
Failure to ventilate
Heart-Lung (Cardio-Pulmonary) Bypass

Switch from anesthesia machine ventilator to heart-lung bypass machine and back.
Heart-Lung (Cardio-Pulmonary) Bypass


“... the anesthesiologist forgot to resume ventilation after separation from cardiopulmonary bypass. The delayed detection ... was attributed to the fact that the audible alarms ... had been disabled during bypass... patient sustained permanent brain damage.”

Almost every surgical team has experienced this error!
Heart-Lung (Cardio-Pulmonary) Bypass

Smart alarm system would provide warning if both ventilator and bypass pump are off.

Should alarm if both are off

NOT Commercially available

Smart alarm system would provide warning if both ventilator and bypass pump are off.
Needs and complexity will only increase …

Point-of-Care Medical Devices (wired ⇒ wireless and mobile)

Data Integration, Analysis, and Display

Connectivity challenge extends beyond the OR

Credit: P. Carleton, RN
Unmet Clinical Needs:
many years of “great ideas” – no implementation pathway

• “Integration of operating room monitors for
development of a smart alarm
system” (Navabi/Mylrea 1990)
• “A system for optimized design of fluid
resuscitation in trauma” (1991)
• OR: 70% of anesthesiologists disable clinical
alarms (Block, Nuutinen, Ballast 1995)
• ICU: 86% false alarms (Tsien, Fackler CCM 1997)
Medical device integration demonstrated at national medical conference
NSF CPS-funded Collaboration: “Architecture for the Safe Composition of Complex Medical Systems”

Goal: Provide technology foundation for real time safety critical systems in healthcare

Specific Issue:
– Real-time safety analysis as multiple devices are integrated
NSF CPS-funded collaboration: “Composition, Interoperability, and Robustness of CPS Components for MD PnP”

- Complex medical device hardware and software requires new methods to certify software prior to clinical use.

- New design and productivity technologies are needed
“a computer interface problem between the CT scanner and the workstation containing the Variseed software occurred that prevented the precise calculation of doses of radiation … received by the patient”*

Dr. Gary Kao’s statement to U.S. Senate Committee on Veterans’ Affairs, June 29, 2009, regarding problems with prostate cancer radiation seed treatments in 92 veterans at Philadelphia VA Medical Center:

*Section 11c and 12e
Medical Device “Plug-and-Play”
Interoperability Program (MD PnP)

Massachusetts General Hospital and the CIMIT, with Army/TATRC support, initiated the MD PnP program in 2004 to lead the adoption of open standards and technology for medical device interoperability to improve patient safety.

More than 85 companies and institutions and > 700 experts (clinicians and engineers) have participated.
Voice of the MD PnP stakeholder community:
Interoperability requires many elements to be aligned

• Focus on **clinical needs**
• **Regulatory** obstacles (*Lui Sha, UIUC*)
• **Liability concerns** (*Insup Lee, Penn*)
• **Business case** (*MD FIRE – Kaiser, Hopkins, Partners HealthCare*)
• Promote/develop/adopt suitable **standards** (*ASTM ICE*)

*Non trivial technology challenges exist in each area*
Goals of the MD PnP Program

1. Lead the adoption of open standards and related technology to support medical device interoperability and system solutions
2. Define a regulatory pathway in partnership with the FDA
3. Elicit clinical requirements for the proposed interoperable solutions
4. Use our vendor-neutral laboratory to:
   • evaluate interoperability standards and solutions
   • serve as a national resource
5. Investigate safety of proposed engineering solutions
New ASTM standard for the **Integrated Clinical Environment**
For safe “plug-and-play” integration of medical devices
• April 2009: The 2nd Joint Workshop On High Confidence Medical Devices, Software, and Systems (HCMDSS) and Medical Device Plug-and-Play (MD PnP) Interoperability.

• 18 submitted papers - specific ICE standard-related implementation research

http://www.cpsweek.org/
Plug-and-Play Network Devices

Another enabling technology for the aforementioned vision is the development of plug-and-play networking technology for medical devices. Plug-and-play capability is needed to ease the setup of integrated point-of-care and extramural arrays of medical devices that communicate with a patient’s electronic health record.

Devising the technology would require addressing concerns about privacy, security, safety, regulations, and technology. In hospital settings, for example, networks would form and reform frequently, as patients are admitted and discharged. Technology for the rapid formation of ad hoc networks needs developing. At the same time, authentication mechanisms would be needed to
MD FIRE

**Medical Device Free Interoperability Requirements for the Enterprise**

- Position Statement & Sample of Interoperability RFP and Contract language
- Developed by Mass General Hospital / Partners, Hopkins, Kaiser
- Conveys healthcare needs to industry, and simplify purchasing specifications
- Released Oct 17, 2008

5 Stakeholder groups from each organization:
Purchasing/materials management, BME, IS, Clinical, Legal

*Download MD FIRE from www.mdapnp.org*
“Healthcare Delivery Organizations (HDOs) must lead a nationwide call to action for interoperability of medical devices and systems. One way that HDOs can effect this change is by including medical device interoperability as an essential element in the procurement process and in vendor selection criteria.”

Download: http://mdpnpc.org/MD_FIRE.php
RESOLVED, That our American Medical Association (AMA) believes that intercommunication and interoperability of electronic medical devices could lead to important advances in patient safety and patient care, and that the standards and protocols to allow such seamless intercommunication should be developed fully with these advances in mind. Our AMA also recognizes that, as in all technological advances, interoperability poses safety and medico-legal challenges as well ...”

as of July 2009:

Anesthesia Patient Safety Foundation
Society for Technology in Anesthesia
Society of American Gastrointestinal Endoscopic Surgeons

American Medical Association
World Federation of Societies of Anesthesiologists
American Society of Anesthesiologists
Massachusetts Medical Society
Financial Implications of standards-compliant medical device interoperability

- Kaiser Permanente: 2006 Analysis medical device-to-EMR integration costs with and without interoperability standards
- Devices: bedside monitors and ventilators
- Analysis excludes safety and workflow benefits
- Results: standard interfaces would reduce integration costs 30%
- Savings: ~ $12M annually
“Automation” in healthcare

• Automation has increased safety and efficiency in aviation, agriculture, manufacturing and some areas of healthcare (e.g. pharmacy).
• Example: Clinical workflow automation, dynamic smart checklists
• System integration and interoperability are necessary for automation

Attribution: L. Sha
Will we reach the tipping point?

- Clinical Push (Societies)
- Hospital Demand (MD FIRE)
- Technology / Platform
- Standards (ICE and others)
- Regulatory (FDA)
- Document Clinical Need
- Alignment with Federal initiatives

Interoperability

Adoption
Our Vision

Improve safety and efficiency by changing expectations; changing technology; changing healthcare.
Adoption of medical device interoperability (standards and technologies) will support:

1. Complete, accurate electronic medical records
2. Rapid deployment of devices in makeshift emergency care settings
3. Clinical decision support systems and smart clinical alarms
4. Support of remote healthcare delivery
5. “flight data recorder” to facilitate adverse events analysis
6. Automated system readiness assessment (prior to starting invasive clinical procedures or critical care transport)
7. Reduce cost of devices and their integration, and reduce EMR-adoptions costs
8. Closed-loop control of therapeutic devices and safety interlocks (e.g. ventilation, medication and fluid delivery)
9. Pathway for innovative medical applications
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