Collaborations

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Learning Objective

Understand how being involved with AAPM and other organizations can also help promote innovative, safe and effective medical devices

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FDA Outreach & Collaboration

- Share information
- Seek input
- Promote collaborations on specific projects

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Food, Drug & Cosmetic Act

- Coordination, research, and outreach are in the law…
  - “plan, conduct, coordinate, and support research, development, training, and operational activities to minimize the emissions of and the exposure of people to, unnecessary electronic product radiation” (360(h)(2))
  - “maintain liaison with and receive information from other Federal and State departments and agencies with related interests, professional organizations, industry, industry and labor associations, and other organizations on present and future potential electronic product radiation” (360(h)(3)).

Collaborations

- The majority of the employees in our divisions are involved in collaborations with federal, state, national, and international groups and individuals across the product areas that we regulate

Federal and State Agencies

- Exchange product safety information
- Exchange information on regulatory activities
- Participate on grant reviews

<table>
<thead>
<tr>
<th>Agency</th>
<th>Description</th>
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<tbody>
<tr>
<td>BARDA</td>
<td>Biomedical Advanced Research and Development Authority</td>
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<tr>
<td>FAA</td>
<td>Federal Aviation Administration</td>
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<tr>
<td>FCC</td>
<td>Federal Communications Commission</td>
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<tr>
<td>CPSC</td>
<td>Consumer Product Safety Commission</td>
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<td>CREPD</td>
<td>Conference of Radiation Control Program Directors</td>
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<td>ESCORS</td>
<td>Emergency Steering Committee on Radiation Standards (ESCOR)</td>
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<tr>
<td>HHS</td>
<td>Health and Human Services</td>
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<td>NIH</td>
<td>National Institute of Health</td>
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<tr>
<td>NIST</td>
<td>National Institute of Standards and Technology</td>
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<tr>
<td>NRC</td>
<td>Nuclear Regulatory Commission</td>
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National Organizations

• Product specific workgroups
• Conference attendance / presentations to learn and share information

<table>
<thead>
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<tbody>
<tr>
<td>AAPM</td>
<td>American Association of Physicists in Medicine</td>
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<td>ACR</td>
<td>American College of Radiology</td>
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<tr>
<td>ASTRO</td>
<td>American Society for Radiation Oncology</td>
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<tr>
<td>Image Gently</td>
<td>Alliance for Radiation Safety in Pediatric Imaging</td>
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<tr>
<td>NCRP</td>
<td>National Council on Radiation Protection and Measurements</td>
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<tr>
<td>QI BA</td>
<td>Quantitative Imaging Biomarker Alliance</td>
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International Organizations

• Share information on radiation safety and regulations
• Develop radiation safety publications

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<tr>
<td>HERCA</td>
<td>Heads of the European Radiological protection Competent Authorities (HERCA)</td>
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<tr>
<td>IAEA</td>
<td>International Atomic Energy Agency</td>
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<tr>
<td>ICRP</td>
<td>International Commission on Radiological Health (ICRP)</td>
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<tr>
<td>WHO</td>
<td>World Health Organization</td>
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Standards and Trade Associations

• Work on product specific standards
• Single point-of-contact to receive/share information with product manufacturers

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<td>AdvaMed</td>
<td>Advancing Medical Technology Association</td>
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<tr>
<td>IEC</td>
<td>International Electrotechnical Commission</td>
</tr>
<tr>
<td>MITA</td>
<td>Medical Imaging Technology Alliance</td>
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Discuss examples related to...

- AAPM Committees and IEC Standards
- Radiation/Product Safety
- Research and Development

(note: in the examples, I have underlined how you can be involved, or may be directly impacted)

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AAPM Committees

- FDA attendees coordinate to cover most AAPM committees, and FDA staff chair/co-chair multiple committees
- Government and Regulatory Affairs (GRAC)
- Technology Assessment Committee (TAC)
- Imaging Physics Committee (IPC)
  - Computed Tomography Subcommittee
  - TG 200 - CT Dosimetry Phantoms and the implementation of AAPM Report Number 111
  - TG 233 - Performance Evaluation of Computed Tomography Systems
- Computer Aided Detection in Diagnostic Imaging (CAD) Subcommittee
- Mammography Subcommittee
- Radiography and Fluoroscopy Subcommittee
  - TG 150 - Acceptance Testing and Quality Control of Digital Radiographic Imaging Systems

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AAPM Committees

- Pediatric Imaging Subcommittee
  - TG 251 - Survey of Pediatric Fluoroscopic Exposure Rates
  - TG 252 - Methodology for the development of pediatric techniques for computed radiography and digital radiography
  - TG 361 - Quality control methodology for low-dose dental and maxillofacial CBCT systems
- Tomosynthesis Subcommittee
  - TG 245 - Tomosynthesis Quality Control
- Work Group on Medical Display
  - TG 196 - Requirements and methods for color displays in medicine
  - TG 260 - Considerations for the Use of Handheld Devices for Viewing Medical Images
  - TG 270 - Display QA
- Working Group on Magnetic Resonance Safety
  - and radiation therapy committees...
International Standards

- Division wants to increase reliance on voluntary consensus standards and issue guidance documents that reference IEC standards or allow conformance to voluntary standards, in lieu of parts of FDA standards

- FDA is engaged with IEC and industry organizations to introduce safety features into the national and international standards for medical (e.g., x-ray, MRI) and non-medical (e.g., laser, microwave) products

- You can be involved in standards development, or related workgroups

IEC Standards: X-Ray Equipment

- FDA is an active participant in the development of CT, fluoroscopy, and radiography IEC standards

- FDA proposals for updates to standards based on AAPM, ACR, Image Gently and input from other professional organizations include requirements to provide QC instructions to users, structured dose reporting, and pediatric safety features

- FDA is working with CRCPD's H-46 Task Force on IEC Standards to develop a pathway to address the practical implementation issues related to performance testing done by FDA, states, and medical physicists

- FDA is seeking to engage you (regulators, industry, and physicists) to explore how to support performance testing to voluntary consensus standards

Radiation Safety

- FDA is actively engaged with international organizations (e.g., WHO, IAEA, HERCA) and researchers on issues related to product and radiation safety

- Recent publications:
  - WHO publication “Communicating radiation risks in paediatric imaging” in April 2016
  - Cancer risks in U.S. radiologic technologists working with fluoroscopically-guided interventional procedures (AJR: Amer J Roentgenol May 2016)
  - Long-term Mortality in 43,763 U.S. Radiologists Compared with 64,990 U.S. Psychiatrists (Radiology July 2016)
  - Organ doses from diagnostic medical radiography – trends over eight decades (1930 to 2010) (Health Physics 2016, in press)
Nationwide Evaluation of X-ray Trends Surveys

- Collaboration among FDA, CRCPD, and ACR
  - Captures radiation exposure data using phantoms from a nationally representative sample of U.S. clinical facilities for specific examinations
  - Publishes the state of the practice of diagnostic radiology
    - November 2015 – Publication of CT Survey (dosimetry for 17 adult and pediatric exams)
    - June 2016 – Summary publication for Cardiac Catheterization Fluoroscopy
    - Upcoming – Dental Radiography (currently analyzing collected data)
- You can help with data collection and analysis

Magnetic Resonance Imaging

- MR Burn Prevention Poster (October 2015)
- Collaboration with members of the International Society for Magnetic Resonance in Medicine
- Incorporates knowledge from review of most frequent adverse events (submitted by you as a facility/manufacturer)
- You can download/print a copy

Radiation Oncology

- Exchange information with ASTRO, NRC, and federal agencies related to adverse events and device failures
- FDA Participation in IEC Subcommittee 62C Radiation Therapy:
  - Discussed key changes in the 4th edition of LINAC standard, IEC 60601-2-1: Particular requirements for the basic safety and essential performance of electron accelerators in the range 1 MeV to 50 MeV
  - Discussion of amendment and new editions of IEC 61217 (Coordinates & Scales) 62083 (Treatment planning systems), 62274 (Record & Verify) and 61168 (Simulator performance)
- Your participation and reporting of adverse event is important to identifying trends and enhance safety standards
FDA Guidance Documents

- Provide another means for stakeholder input, and consistency
  - FDA’s Guidance for the submission of 510(k)s for Solid State X-ray Imaging Device recommends a list of tests to be performed for digital radiographic systems with solid state x-ray imagers in premarket submissions (http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/ucm073780.htm)
  - TG150 (Acceptance Testing and Quality Control of Digital Radiographic Imaging Systems) is currently working on a AAPM report for a list of recommended acceptance and routine tests for digital radiographic systems
- Your collaboration/comments promote use of the latest science and consistency in premarket testing and clinical quality control

FDA Public Advisory Panel Meetings

- FDA holds public advisory panel meetings to seek feedback from panel experts, patients, members of the public, and regulated industry
- You can apply to be a member of the Radiological Advisory Committee (http://www.fda.gov/AdvisoryCommittees/default.htm)
- You can present as a member of the public at these meetings, or send comments regarding the meeting materials and/or draft guidance documents

FDA Communications

- You can sign-up to you to receive important FDA news and information as it become available (e.g., Recalls and Safety Alerts, Medical Devices, Radiological Health, FDA Press Releases)
- Note: options for medical and non-medical radiation emitting electronic products that FDA regulates (e.g., lasers, microwaves, sunlamps)
- FDA also publishes notices of guidance documents, upcoming meetings, and proposed regulations in the Federal Register

http://www.fda.gov/AboutFDA/ContactFDA/StayInformed/GetEmailUpdates/default.htm?source=govdelivery
Product Development

- FDA’s mission to protect and promote public health depends on safe and effective products coming to the market

Pre-Submissions

- FDA/CDRH’s pre-submission program allows you (researchers or manufacturers) to request feedback from the FDA on proposed regulatory pathway and protocols
  - (also for study risk determination, information, and submission issue meetings)
- Consider for devices with novel or specific intended uses

Grant Reviews

- Our division occasionally receives a request to provide feedback on the application and/or regulatory pathway discussion for grant proposals for innovative therapeutic and diagnostic devices
  - Examples: BARDA and NIH grants
- Our feedback provides grant applicants with a better understanding the regulatory pathway for new products

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Quantitative Imaging Biomarkers Alliance

- Collaboration to create consistent, reliable, valid and achievable quantitative imaging results across imaging platforms, clinical sites, and time.
- Accelerating development and adoption of hardware and software standards to achieve accurate and reproducible quantitative results from imaging methods.
- FDA is well represented across the committees and has lead related research projects.
- You can become involved to support the research and development.

https://www.rsna.org/qiba/

Research Collaborations

- Supports the evaluation of safety and effectiveness of products:
  - Phantom development for breast imaging.
  - Computed aided diagnosis and methodologies for algorithm assessment:
    - Computer Aided Detection in Diagnostic Imaging (CAD) Subcommittee (Berkman Sahiner, SC Chair).
  - Virtual imaging clinical trials for regulatory evaluation (VICTRE):
  - CT iterative reconstruction – object assessment of image quality and dose reduction.

- You can collaborate, or download imaging system simulation tools and statistical analysis software for reader or algorithm assessment that CDRH makes public.

https://github.com/DIDSR

Summary

- FDA engages in multiple efforts to seek public input, and collaborate with federal, state, national, international groups, and individuals.
- Being involved with AAPM and other organizations and sharing your research/physics/clinical expertise helps promote safe and effective medical devices.

Contact Information and Resources

Division of Radiological Health
Robert.Ochs@fda.hhs.gov

Device Advice: Comprehensive Regulatory Assistance
http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/default.htm

CDRH-Division of Industry and Consumer Education (DICE)
DICE@fda.hhs.gov

1. Pre-market
2. Post-market
3. Regulatory research
4. Collaborations