



IMDRF

International Medical
Device Regulators Forum

STANDARDS WORKING GROUP UPDATE





STANDARDS WORKING GROUP (SWG)

- **NWIP Goal**
 - Improve the utility of standards for regulatory use in order to streamline review processes and harmonize regional and national regulatory approaches
- **Objectives**
 1. Background research:
 - Identify problems in standards development that diminish their regulatory utility
 - Analyze IMDRF member engagement with Standards Developing Organizations (SDOs)
 2. Draft recommendations for developing 'regulatory-ready' standards
 3. Enhance IMDRF relationships with ISO and IEC



NWIP OUTCOMES

- 2017 report to Management Committee
 - *Improving the Quality of International Medical Device Standards for Regulatory Use*
- 2018 draft guidance for public consultation
 - *Optimizing Standards for Regulatory Use*
- Strong and growing relationships with ISO and IEC
 - Agreement with IEC
 - Liaison A status with ISO TC210 pending ISO resolution



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OUTCOME: MC REPORT

- **Audience**
 - Management Committee members
 - IMDRF members
- **Background research**
 - Many standards not useful for regulators
 - Regulatory Authorities' (RAs') participation in ISO and IEC is inconsistent, at both national and international levels
 - Standards created with regulatory purposes in mind can streamline and harmonize regulatory processes
- **Proceedings from ISO/IEC/IMDRG SWG workshop**
 - SDOs welcome greater regulator and IMDRF engagement



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OUTCOME: DRAFT GUIDANCE

- **Audience**
 - Regulatory Authorities
 - SDOs
 - Stakeholders interested in standards' improvement for regulatory purposes
- **Recommendations**
 - For standards development
 - For participation in ISO and IEC
 - For future IMDRF engagement



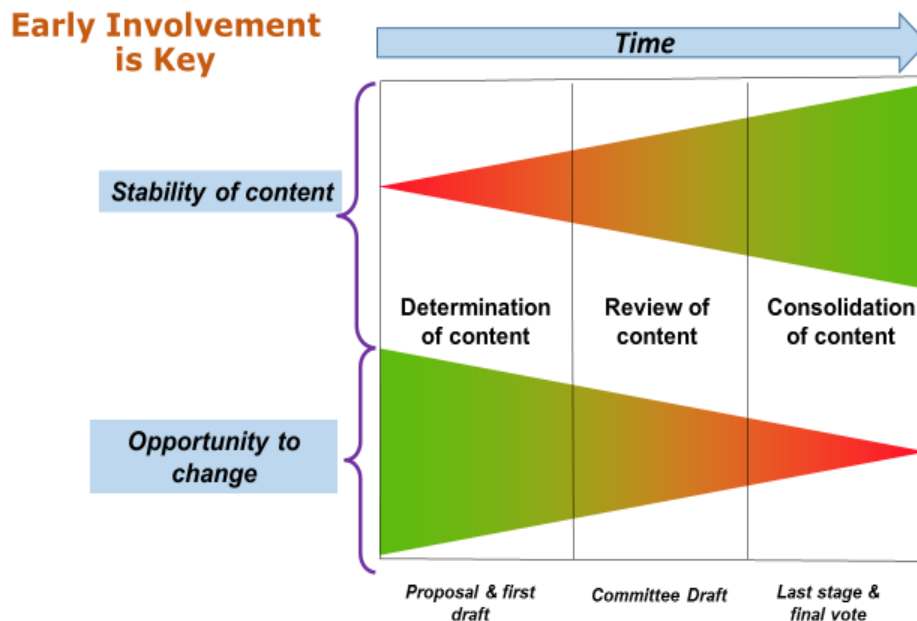
GUIDANCE: STANDARDS DEVELOPMENT

- **Optimizing standards' content, e.g.,**
 - Elements for inclusion
 - Attention to appropriate rationale
 - Straightforward and clear conformance acceptance criteria
- **Best practices for standards procedures, e.g.,**
 - Applying consensus principles
 - Emphasis on RAs' contributions
 - Transparency on authorship of standard and comments



GUIDANCE: RA PARTICIPATION

- **Engagement: why and how to work with**
 - National Bodies and mirror committees
 - SDOs at the international level
- **Effective commenting: quality and timing**





GUIDANCE: IMDRF ENGAGEMENT

- IMDRF enjoys a unique position of authority in device regulation harmonization
- IMDRF standards group offers opportunity for RAs to speak with one voice to SDOs
- IMDRF can
 - Act as a resource and communications hub to both members and SDOs
 - Advance regulatory science



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OUTCOME: SDO RELATIONSHIPS

- **ISO**

- IMDRF has formally applied to TC210 for Category A liaison status
- Resolution at Technical Committee level is required; Chair is SWG member

- **IEC**

- IMDRF has formally applied to TC62 and its subcommittees for Category A liaison status
- Memo of Understanding under review



NEXT STEPS

- **Short term**

- Public consultation of the IMDRF draft guidance *Optimizing Standards for Regulatory Use* closes 24 May 2018
- Standards working group meeting in June 2018 to resolve comments and plan for NWIP ‘
- Finalize guidance by Sept 2018, then promote and educate

- **Medium term**

- Advance SDO relationships/agreements
- Discern how to effectively represent IMDRF members in standards development priorities
- Operationalize liaison status and MoU/agreements with SDOs



NEXT STEPS (CONT'D)

- **Longer term – consider sustainability**
 - Analyze further how standards' can contribute to IMDRF strategic goal to '...accelerate international medical device regulatory convergence...'
 - Determine appropriate future role for standards in IMDRF
 - Liaise with SDOs
 - Lead productive participation in standards development ('voice of regulators')
 - *Drive application of standards to regulatory convergence – how can we put standards to work on behalf of harmonization?*



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NEW WORK ITEM PROPOSAL

Recognition of International Standards by IMDRF members

- Update the list of international standards recognized by IMDRF members.
- Analyze the differences and similarities in adoption/recognition policies and processes among members.
- Research on technical difference in adoption/recognition of these standards among members.
- Develop a guidance for IMDRF members with recommendations on the effective use of a recognition program to advance regulatory science and streamline regulatory activities



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THANK YOU