Regulatory Submissions, Information, and Document Management Forum

FEBRUARY 14-16, 2022 | BETHESDA NORTH MARRIOTT HOTEL AND CONFERENCE CENTER | NORTH BETHESDA, MD



CALL FOR ABSTRACTS

PRESENTATION ABSTRACT SUBMISSION DETAILS AND GUIDELINES SUBMISSION DEADLINE: AUGUST 10

Are you a Professional involved in regulatory informatics and data, intelligence, electronic submissions, or document management? If so, DIA wants to hear from YOU! The *Regulatory Submissions, Information, and Document Management (RSIDM) Forum* will cover topics crossing all areas of regulatory information management, separated into four tracks: Regulatory Informatics for Business and Technology, Electronic Document Management (EDM), and Electronic Regulatory Submissions (ERS). Please select the track that best fits with your proposal. If your topic is relevant to more than one track, please indicate that in your abstract summary as cross-track sessions are available. Both in-person and virtual presentations will be considered. Please indicate whether you will anticipate presenting in-person at the hotel or if you will be remote.

The RSIDM Program Committee is seeking proposals for Presentations, Sessions, and Short Courses on the following topics (keep in mind, business use cases and lessons learned are encouraged in all topic areas). As this forum is attended by many regulatory professionals, service providers and health authority representatives, topics in addition to those listed below that you feel are relevant may be submitted for evaluation and possible selection.

Regulatory Informatics: NEW FOCUS (formally known as Regulatory Information Management (RIM))

The Regulatory Informatics track provides the forum for discussions, information sharing, and best practices on managing regulatory data across the product development lifecycle for pharmaceuticals, medical devices, diagnostics, consumer products, and more from R&D to post-marketing activities within organizations and health authorities and the broader healthcare ecosystem.

- · Business Process Implications of Regulatory Data
- RIM Implementation Case Studies and Best Practices
- Data Strategy and (Master) Data Management: Use of technology, applications, methodologies, and operating models
- Data Quality: Measuring, Sustaining, Remediating, and Emerging Technologies (e.g., Artificial Intelligence [AI]/Machine Learning [ML])
- Data Standards: Interoperability, Reference Models, and Regulatory Requirements (e.g., IDMP, PQ-CMC, ePI)
- Data Governance: Methodologies, Technologies, Operating Models, and Data Governance Programs
- Planning, Tracking and Management of Regulatory Activities and Information
- Regulatory Analytics: Reporting, Operational KPIs, Metrics and Insights for Forecasting and Strategic Decision Making
- · Organizational Change Management
- Regulatory Intelligence and Regulatory Strategy
- Cross-functional Dependencies on Regulatory Data (e.g., Clinical, Supply Chain, Manufacturing, Safety, and Commercial)
- Digital Transformation of Regulatory through Intelligent Automation (e.g., RPA/AI/ML)
- Regulatory Capabilities to Support Medical Devices and Diagnostics: RIM, Software as a Medical Device
- Impact of Mergers/Acquisitions and Product Divestitures and Product Divestitures on Regulatory Information
- Partnerships/Outsourcing
- Health Authority, Industry and Trade Group Vision and Collaborations
- Structured Content Management

Electronic Regulatory Submissions (ERS)

This track explores the submission process, regulatory requirements and new developments, best practices and case studies in regulatory submissions and industry adoption techniques.

- · Automation and Artificial Intelligence
- · Dossier Lifecycle Management
- Global Filings Strategies (e.g., Agency Pilots)
- · Validation/Quality Strategies
- Specialty Submissions (e.g., AdPromo or Establishment Registration)
- · Portfolio and Global Submission Management
- Operational Efficiencies and Reuse
- Planning, Tracking, and Metrics

- · Mergers/Acquisitions and Product Divestitures
- Partnerships: Contracting Considerations, Content Collaboration and Re-Use, Shared systems
- Outsourcing: Vendor Selection, Oversight, and Relationship Management
- Topics Relative to Small Pharmaceutical Companies and Organizations
- Building a Reg Ops Team: Expanding Reg Ops Infrastructure and Skill Sets, Training, and Managing Increasing Demands
- · Future of Submissions: Documents to Data
- Global Submission Specifications
 - EMA Clinical Trials Information System CTIS impacts
 - Brexit
 - · DADI and Application Forms
 - eCTD 4 Pilot Activities

Electronic Document Management (EDM)

This track examines the processes, systems, and best practices for content management across the product lifecycle, including alignment with the RIM system for optimal use of regulatory information.

- Intelligent Automation (e.g., using bots and natural language processing) and Working with Validation Content Management Systems - Case Studies/ Experiences
- Trial Master File (TMF) and eTMF Operations
- Optimizing the Clinical/ Regulatory/ Quality Common Content and Processes
- Collaboration Across the Organization and with CROs, Partners, and Vendors
- Regulatory Submission Documents/eCTD
- Inspection Preparation and Readiness
- · Quality Management Systems and Quality Control
- Labeling Lifecycle
- Structured Authoring and Workflows: Case Studies/Experiences
- Effective Change Control Management (e.g., Labeling, CMC, Supply Chain)
- Integration Between Regulatory Informatics and EDM Systems; Business Process, and Across Informatics and Document Management Capabilities (e.g., RIM and Regulatory Content; eTMF and CTMS)
- Mergers/Acquisitions and Product Divestitures

Abstract Submission Deadline: Friday, August 10

Notification: Week of September 20

Final PowerPoint Presentations Due: January 24, 2022

Forum Dates: February 14-16, 2022

Please submit all abstracts online at: <u>DIAglobal.org/Abstracts</u>

Questions: Contact Jessica L. Roman, MS, CMP, Senior Project Manager, at <u>Jessica.Roman@DIAglobal.org</u>

CALL FOR PRESENTATION ABSTRACTS

SUBMISSION DEADLINE: AUGUST 10

GENERAL SUBMISSION REQUIREMENTS

- · All submissions must be submitted online
- For complete submission requirements and to submit your abstract go to DIAglobal.org/Abstracts

SUBMISSION TIPS

- Ideal submissions will contain practical content and shared experiences
- Theoretical topics and content is acceptable, however, it should be supported with proof of concepts and use cases
- Diverse topics and sessions are welcomed and encouraged within the scope of the forum
- Please select the track that best fits with your proposal. If your topic is relevant to more than one track, please indicate that in your abstract summary as cross-track sessions are available.
- Abstracts should be written using clear language and descriptions to provide enough clarity for the selection committee to review and understand

REQUIRED DOCUMENTATION FOR ALL ABSTRACTS

- Participant Disclosure Information: All abstract authors must disclose any
 relevant financial relationships with any commercial interest associated with
 this activity that exist or have existed within the past 12 months, as well
 as any discussion of unlabeled or unapproved drugs or devices. If you are
 proposing an abstract on behalf of the author, as the submitter you will not
 be asked to disclose. However, should the abstract be accepted, the author
 will be informed that he or she must complete and submit a Participant
 Disclosure in order to participate in the program
- All submitters and authors must agree to the DIA Speaker Authorization for Use of Presentation Materials in order for the abstract to be a part of the Program. Accepted abstracts will be available on DIA's website for attendee download.

SUBMISSION GUIDELINES

Submitting a PRESENTATION ABSTRACT (All abstracts must be submitted online)

15-20-minute presentation, bundled with other presentations to create a session. Abstract author is considered the presenter (co-presenters are not permitted) and will be responsible for:

- Adhering to the program development guidelines and timelines
- Working with chair and other presenters in creating a balanced program offering
- Preparing and delivering a PowerPoint presentation

Submitting a SESSION ABSTRACT (All abstracts must be submitted online)

75-minute total session. Abstract author will be responsible for:

- · Adhering to the program development guidelines and timelines
- Recruiting speakers and ensuring good representation/diversity in their selection. Maximum of 3 speakers per session
- Working with the Session Chair to communicate with speakers regarding their role in the session

Submitting a SHORT COURSE ABSTRACT (All abstracts must be submitted online)

Three-hour, interactive presentation delivered in small group format. Abstract author is considered the Short Course Lead Instructor and will be responsible for:

- · Adhering to the program development guidelines and timelines
- Recruiting co-instructors and ensuring good representation/diversity in their selection
- Communicating with co-instructors regarding their role in the short course and reviewing presentation materials (note: PowerPoint presentations are required from each instructor)
- Managing the short course, including the facilitation of audience questions and interactions

SUBMISSION GUIDELINES

The following information will be requested at the time of submission. **DIAglobal.org/Abstracts**

To streamline your submission process and avoid possible delays, DIA strongly encourages you to submit your abstract as early as possible. **Do not wait until the last day.**

Prepare your abstract in advance of accessing the DIA website. Abstract information should be copied and pasted from a prepared document as plain text. **All of the below fields are required.**

Submitter or Author Information

Prefix: Country:

First Name: Address Line:

Middle Name: City:

Last Name: State/Province:

Name Suffix: Zip/Postal Code:

Email:

Degrees: Phone:

Job Title: Company:

NOTE: If you are submitting on behalf of author, you are considered the SUBMITTER and will need to complete the required information for yourself and also for the AUTHOR. Submitters will be the contact for author regarding the status of the abstract.