

# eRegulatory Summit

8 - 10 April 2019  
Crowne Plaza Fira Center  
Barcelona, Spain

## MEET THE INNOVATORS CHANGING THE FACE OF INFORMATION MANAGEMENT AND ESUBMISSIONS

Telematics, RIM, eCTD, IDMP  
and Filing Variations:  
Actionable insights to improve  
your regulatory strategy



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# Day 1: Monday 8th April

	Global Submissions	Telematics & RIM
08.00	<i>Registration</i>	
09.00	<b>Opening Remarks</b>	<b>Opening Remarks</b>
09.10	<b>CASE STUDY: Implementing eCTD within National Procedures</b> <ul style="list-style-type: none"> <li>• Pain points from industry during the implementation of eCTD in National Procedures</li> <li>• Understanding the value in creating a baseline submission</li> <li>• Best practice for harmonising dossiers</li> </ul> <b>Daniel Verrall</b> , Senior Team Manager, <b>Pfizer</b>	<b>Feedback on EMA's IDMP Implementation Guidelines</b> <ul style="list-style-type: none"> <li>• Reviewing EMA's implementation guide, the defined items and timelines</li> <li>• Understanding the importance of the IDMP Implementation Guidelines for successful transition to IDMP</li> <li>• Designing a strategic implementation plan</li> </ul> <b>Laurent Desqueper</b> , XEVMPD & IDMP Business System Owner, <b>MSD Europe Inc.</b>
09.45	<b>Regulatory Agency Perspectives: Feedback, plans and practical advice for eCTD submissions moving forward</b> <ul style="list-style-type: none"> <li>• Exploring current regulatory timelines</li> <li>• Discussing common pitfalls during eSubmissions process and how they can be avoided</li> <li>• Examples of handling industry data submissions</li> </ul> <b>Jaana Pohjonen</b> , Specialist of Record Management and Archives, Quality Manager, <b>FIMEA</b>	<b>Lessons Learned from Data Migration</b> <ul style="list-style-type: none"> <li>• Case study: end-to-end execution of data migration</li> <li>• How best to maintain data quality during migration?</li> <li>• Common problems faced during the migration process</li> </ul> <b>Danielle Beaulieu</b> , Director, Global Regulatory Business Capacities, <b>Bristol-Myers Squibb</b>
10:20	<b>An Update on Clinical Trial Regulation in the EU</b> <ul style="list-style-type: none"> <li>• Current and projected timelines for the EU CTA</li> <li>• Working through the Clinical Trial Regulation and submissions portals</li> <li>• Discussing best practice for the incorporation of clinical trial submissions into eCTD</li> </ul> <b>Speaker to be Announced</b>	<b>Improving Data Integration Across the Business</b> <ul style="list-style-type: none"> <li>• Discussing the significance of information integration</li> <li>• Data integration implementation best practice</li> <li>• Current challenges in cooperation across departments</li> <li>• Leveraging cross-functional expertise to strengthen relationships</li> </ul> <b>Marty Boom</b> , Global Head of Regulatory and Safety, <b>Navitas Life Sciences</b>
10.55	<i>Morning Coffee &amp; Networking</i>	
11.20	<b>Best Practice for Implementing the EU Falsified Medicines Directive</b> <ul style="list-style-type: none"> <li>• EU-Falsified Medicines Directive What is required? Timeline and implications</li> <li>• How to implement the unique identifier and the tamper verification feature?</li> <li>• Effects and consequences for the pharmaceutical and packaging industry</li> </ul> <b>Johan Verhaeghe</b> , National Policy Liaison, <b>Medicines for Europe</b>	<b>Data Modelling for Regulatory Affairs</b> <ul style="list-style-type: none"> <li>• Exactly know what you have</li> <li>• Understand the end to end relationship within the lifecycle of those data</li> <li>• Avoid data redundancies</li> <li>• Improve data quality</li> <li>• Form the prerequisite for any proper digitalization activities.</li> </ul> <b>Dr. Jörg Stüben</b> , Head of Regulatory Information Management and Senior Expert, <b>Boehringer Ingelheim International GmbH</b>
11.55	<b>Experiences of Submitting Dossiers in China</b> <ul style="list-style-type: none"> <li>• Industry experience of submitting in China</li> <li>• Reflection on the changes in the regulatory environment</li> <li>• Updates on eCTD adoption status and trends</li> </ul> <b>Speaker to be Announced</b>	<b>Utilizing AI and Data Analytics to Optimise your Regulatory Strategy</b> <ul style="list-style-type: none"> <li>• How you can use AI to get more out of your existing data</li> <li>• Exploring the impact of AI on the regulatory submissions process</li> <li>• Defining and understanding the main challenges for stakeholders</li> </ul> Representative from <b>Cunesoft</b>
12.30	<i>Networking Lunch</i>	

	Global Submissions	Telematics & RIM
14.00	<p><b>Assessing the Latest Developments Towards eCTD in Singapore</b></p> <ul style="list-style-type: none"> <li>• Outlining the current regulatory review process and good review practices</li> <li>• Updates on eCTD adoption in Singapore</li> <li>• Discussing common submission pitfalls</li> </ul> <p><b>Speaker to be Announced</b></p>	<p><b>INDUSTRY CASE STUDY: Cross Functional Collaboration – Data Integration Across the Business</b></p> <ul style="list-style-type: none"> <li>• Discussing the significance of information integration</li> <li>• Data integration implementation best practice</li> <li>• Current challenges in cooperation across departments</li> <li>• Leveraging cross-functional expertise to strengthen relationships</li> </ul> <p><b>Patrick Middag</b>, Associate Director, Regulatory IT, <b>Bristol-Myers Squibb</b>  <b>Deborah McCloskey</b>, Global Regulatory Business Capabilities, <b>Bristol-Myers Squibb</b></p>
14.35	<p><b>Guidance from South East Asia on the Harmonisation and Implementation of eCTD</b></p> <ul style="list-style-type: none"> <li>• Where do we stand with the ASEAN Pharmaceutical Harmonisation Standard?</li> <li>• Discussing the latest eCTD updates and expectations in Thailand</li> <li>• Updates and timelines from around the region</li> </ul> <p><b>Speaker to be Announced</b></p>	<p><b>Industry Feedback on the Falsified Medicines Directive: Post Implementation</b></p> <ul style="list-style-type: none"> <li>• Feedback from industry on the implementation best practice</li> <li>• Pragmatic advice on strategies for departmental data coordination</li> <li>• Case study on minimising risk and data alignment to fill potential data gaps</li> </ul> <p><b>Quentin Grignet</b>, IDMP Project Lead, <b>GSK</b></p>
15:10	<p><b>eSubmissions Strategy at a SME</b></p> <ul style="list-style-type: none"> <li>• Best practice for implementing an efficient eCTD strategy</li> <li>• Deciding factors: in-house or outsourcing? The best options for your regulatory submissions process</li> <li>• Outlining challenges and sharing experience of overcoming them</li> </ul> <p>Representative from <b>Veeva</b></p>	<p><b>Spotlight Session</b></p> <p>Representative from <b>Amplexor</b></p>
15.45	<i>Coffee &amp; Networking Break</i>	
16.15	<p><b>Outlining Current Regulatory Submission Status in Jordan</b></p> <ul style="list-style-type: none"> <li>• Latest updates from the JFDA including new eCTD specification</li> <li>• Outlining the current regulatory review process and good review practices</li> <li>• Discussing common pitfalls during submissions process and how they can be avoided</li> </ul> <p>Representative from <b>Jordan FDA</b></p>	<p><b>Designing an End-to-End RIM System</b></p> <ul style="list-style-type: none"> <li>• Understand the increasing importance and roles of RIM systems within organizations</li> <li>• Formulating a foundational RIM strategy</li> <li>• Operational hurdles for smaller companies</li> </ul> <p><b>Costas Mistrellides</b>, IDMP Regulatory Affairs Coordinator, <b>Johnson &amp; Johnson</b>  <b>Chris Dunn</b>, IDMP Specialist, <b>Johnson &amp; Johnson</b></p>
16.50	<p><b>Updates on Submissions in Australia</b></p> <ul style="list-style-type: none"> <li>• Discussing regulation changes out of the TGA in Australia</li> <li>• Sharing practical experiences of successful submissions</li> <li>• Outlining challenges and sharing experience of overcoming them</li> </ul> <p><b>Henrietta Dehmlow</b>, Head of Submission Management, <b>Roche</b></p>	<p><b>PANEL DISCUSSION: The Future of Information Management - Shifting the Mindset from Tactics to Strategy</b></p> <p>The separation between eSubmissions and Data Management is fairly pronounced now, but in a few years' time we will see the two areas intersect. What is the 5-year plan? Where do we see the industry going?</p> <p><b>Joerg Stueben</b>, Senior Expert in Global Regulatory Operations, <b>Boehringer Ingelheim International GmbH</b>  <b>Frits Stulp</b>, IDMP Program Manager, <b>Astellas</b></p>
17:20	<i>Chairman Closing – End of Day 1 and Networking Drinks</i>	

# Day 2: Tuesday 9th April

Plenary Session	
08.00	<i>Registration</i>
09.00	<b>Opening Remarks</b>
09.10	<p style="text-align: center;"><b>Telematics Landscape and Implementation Feedback</b></p> <ul style="list-style-type: none"> <li>• An overview of the current regulatory landscape and projected timelines for implementation                             <ul style="list-style-type: none"> <li>• Feedback on current progress</li> </ul> </li> <li>• Implications for industry: what is expected and how to achieve this</li> </ul> <p style="text-align: center;"><b>Remco Munnik</b>, Regulatory Information Director, <b>Asphalion S.L.</b></p>
09.45	<p style="text-align: center;"><b>Best Practice for Successful Implementation of OMS &amp; RMS</b></p> <ul style="list-style-type: none"> <li>• A detailed look at the operation of OMS &amp; RMS implementation processes                             <ul style="list-style-type: none"> <li>• Discussing the triumphs and hurdles for successful implementation</li> </ul> </li> <li>• Integration of SPOR into the Application Dataset Management module in CESSP</li> </ul> <p style="text-align: center;"><b>Dominik Gigli</b>, Senior Manager IDMP Office, <b>Merck KGaA</b></p>
10:20	<p style="text-align: center;"><b>A Systematic Approach to PMS Preparations</b></p> <ul style="list-style-type: none"> <li>• What is the current outlook for product implementation?</li> <li>• Hit the ground running: discussing techniques and methodology for successful integration in the first instance                             <ul style="list-style-type: none"> <li>• Potential challenges and how best to avoid them</li> </ul> </li> </ul> <p style="text-align: center;"><b>Kelly Hnat</b>, Principal, <b>K2 Consulting</b></p>
10.55	<i>Morning Coffee &amp; Networking</i>
11.25	<p style="text-align: center;"><b>The IDMP Substance Management System</b></p> <ul style="list-style-type: none"> <li>• An update on the FDA's progress with the implementation of G-SRS                             <ul style="list-style-type: none"> <li>• Discussing the status of SMS implementation in the EU</li> </ul> </li> <li>• Examining the differences between systems and the impact of the HMA endorsement of an EU G-SRS project</li> </ul> <p style="text-align: center;"><b>Frits Stulp</b>, IDMP Program Manager, <b>Astellas</b></p>
12.00	<p style="text-align: center;"><b>CESSP Essentials: What to Expect from Phase 1</b></p> <ul style="list-style-type: none"> <li>• Discussing the timelines for submission roll out and implications for the future                             <ul style="list-style-type: none"> <li>• Exploring the interaction of current telematics strategies and CESSP</li> <li>• Preparing a system integration plan to control your master data</li> </ul> </li> </ul> <p style="text-align: center;"><b>Speaker to be Announced</b></p>
12.30	<i>Networking Lunch</i>
Global Submissions	Telematics & RIM
<p>14:00 <b>GCC Experiences with eCTD Transitions and Timelines</b></p> <ul style="list-style-type: none"> <li>• Discussing regulation changes across the GCC region</li> <li>• Sharing practical experiences of successful submissions</li> <li>• Comparing the GCC eCTD submissions process with the EU</li> </ul> <p><b>Richard Knowles</b>, Senior Manager GRAAS Operations, <b>Amgen</b></p>	<p><b>How eCTD and IDMP Will Work Together</b></p> <p>As, strategically, Brexit will impact parts of the telematics strategy (in Europe). My part is mainly looking at how the two worlds should fuse together to form a company's future RIM programme. The presentation will be more strategic as it's making sure customers foresee how their processes will look in the future when IDMP and eCTD do fuse.</p> <p><b>Anjana Pindoria</b>, Director Product Strategy, <b>Extedo</b></p>
<p>14:35 <b>Outlining the Latest Updates from Canada and the Implications for Industry</b></p> <ul style="list-style-type: none"> <li>• Examining the latest results from the eCTD pilot for CTAs</li> <li>• Covering all requirements for 2019</li> <li>• Case study on common submission errors and troubleshooting challenges</li> </ul> <p><b>Susanne Picard</b>, President/General Manager, <b>SPharm</b></p>	<p><b>Label and Artwork Management Systems: Transitions and Sustainability</b></p> <ul style="list-style-type: none"> <li>• How to create a fool-proof plan based on market regulations, products and packaging needs</li> <li>• Best practice for preventing costly delays</li> <li>• Success stories: improving data quality in your tracking database</li> </ul> <p><b>Bas van Heijst</b>, Associate Director Regulatory Affairs, <b>Astellas</b></p>

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## Day 2: Tuesday 9th April *(continued)*

	Global Submissions	Telematics & RIM
15.10	<b>Creating a Global Dossier Plan for Fast-tracking Approvals</b> <ul style="list-style-type: none"> <li>Guidelines and updates on global systems</li> <li>Overview and case studies for expedited approvals</li> <li>Regulatory hurdles and creating a plan to overcome them</li> </ul> <b>Olga Alfieri</b> , Director, Global Submission Management, <b>Eisai</b>	<b>Spotlight Session</b> Representative from <b>ArisGlobal</b>
15:40	<i>Coffee &amp; Networking Break</i>	
16.10	<b>Creating a Common Clinical Trials Submission Format</b> <ul style="list-style-type: none"> <li>Best practice for creating a harmonised Clinical Trial Application format</li> <li>Discussing formats and procedures around the globe</li> <li>Identifying roadblocks and planning for them</li> </ul> <b>Speaker to be Announced</b>	<b>Digital Transformation of Novartis' Regulatory Processes</b> <ul style="list-style-type: none"> <li>Novartis' strategic digital roadmap to transform Regulatory processes</li> <li>Technology enablers</li> <li>Gain operational efficiency through automation</li> <li>Using data insights and predictive analytics to support better decisions</li> <li>Key takeaways</li> </ul> <b>Wolfgang Schleifer</b> , Service Delivery Lead Regulatory, <b>Novartis</b>
16.45	<b>PANEL SESSION: Sharing Experiences of Global eCTD Submissions</b> <ul style="list-style-type: none"> <li>Global eCTD adoption status, trends and v4.0 outlook</li> <li>Sharing experiences of confusion and conflicts from SME and large pharma standpoint</li> <li>Q&amp;A: answering best practice questions</li> </ul>	<b>Developing a Robust Data Archiving System</b> <ul style="list-style-type: none"> <li>Constructing a flexible long-term data archiving system</li> <li>Reviewing options for maintaining data integrity in your archive system</li> <li>Looking at current automation initiatives and how to use them</li> </ul> <b>Michel Mikhail</b> , Expert in International Regulatory Affairs and Biosimilars
17:20	<i>Chairman's Closing Remarks</i>	

## Day 3: Wednesday 10th April

	Filing Variations Focus Day
08.30	<i>Registration</i>
09.00	<b>Speakers Remarks</b> <b>Gry Agapitos</b> , Senior Regulatory Affairs Professional, <b>ALK Abello</b> <b>Justyna Kwiatkowska</b> , Regulatory Affairs Specialist, <b>Adamed Group</b> <b>Monique Mendel Ott</b> , Manager Global Regulatory Affairs, <b>Grünenthal GmbH</b> <b>Sophie Nageotte</b> , Regulatory CMC Consultant
09.10	<b>An Update on ICH Q12</b>
09.45	<b>A Detailed Look at the Variation Classification Guideline</b>
10:20	<b>Demonstrating an Effective Grouping and Work Sharing Strategy</b>
10.55	<i>Morning Coffee &amp; Networking</i>
11.25	<b>Troubleshooting your EU Filing Variation Plan</b>
11.55	<b>Comparing Requirements: Filing Variation Differences between EU and USA</b>
12.35	<b>Outlining the Everchanging Requirements in Asia</b>
13.10	<i>Networking Lunch</i>
14:30	<b>Designing a Successful Global Variation Strategy</b>
15:05	<b>Q&amp;A – Variations Around the World</b>
15:30	<b>Chairman's Closing Remarks</b>