Multi-attendee discounts are available!

FDANEWS PRESENTS THE

EU-MEDICAL DEVICE REGULATION/ IN VITRO DIAGNOSTICS REGULATION COMPLIANCE WORKSHOPS

SEPT. 10-12, 2019 DOUBLETREE BY HILTON PHILADELPHIA CENTER CITY PHILADELPHIA, PA

INTERACTIVE WORKSHOP SERIES FROM FDANEWS AND OMBU ENTERPRISES

Agenda

<u>Day 1</u>

8:00 a.m. – 9:00 a.m.

Registration and Continental Breakfast

- 9:00 a.m. 10:30 a.m.
- Part A Overview of the Regulations
 - Introductions
 - Reasons for Change
 - Structure of the Regulations
 - Notified Bodies
 - Transition Timelines
 - Understanding Standards and Harmonization
 - EU version of ISO 13485:2016 and ISO 14971:2007

Part B – MDR Article 120 Transition

- Status of the EU-MDR Implementation
- Article 120 Transitional Provisions
- Article 120
- ISO 13485:2016 & CEN/TR 17223:2018
- ISO 14971:2019
- Status of NBs
- Harmonized Standards & Common Specifications
- Status of Eudamed
- Manufacturer's Potential Hurdles
- Questions

Part C – Obligations of the Manufacturer

• Economic Operators

	 Single Registration Number QMS, RMS, and PMS CE Mark and Declaration of Conformity UDI requirements Incidents and Field Safety Corrective Actions Person Responsible for Regulatory Compliance Product Liability Insurance Exercise C1 – Person Responsible for Regulatory Compliance Exercise C2 – Economic Operator 	
10:30 a.m. – 10:45 a.m.	Break	
10:45 a.m. – 12:00 p.m.	Part D – The Quality Management System (QMS)	
	 ISO 13485:2016 & CEN/TR 17223:2018 QMS Requirements QMS Certificates Exercise D1 – Elements of the QMS 	
12:00 p.m. – 1:00 p.m.	Lunch Break	
1:00 p.m. – 2:15 p.m.	Part E – The Risk Management System (RMS)	
	 Requirements Description of Changes for ISO 14971:2019 The Process Flow in ISO 14971:2019 Annex I(3) The Risk Management System Annex I(4) Risk Control Measures Annex I(5) Use Error 	
2:15 p.m. – 2:30 p.m.	Break	
2:30 p.m. – 5:00 p.m.	Part F – The Post-market Surveillance System (PMSS)	
	 Risk Management Benefit-Risk Determination (BRD) Clinical Evaluation (CE) Post-market Clinical Follow-up (PMCF) Post-Market Surveillance Post-market Surveillance Report (PMSR) 	

- Periodic Safety Update Report (PSUR)
- Summary of Safety and Clinical Performance (SSCP)
- Incident Reporting
- Exercise F1 Elements of the PMSS

5:00 p.m. And much more!

<u>Day 2</u>

8:00 a.m. – 9:00 a.m.	Registration and Continental Breakfast
9:00 a.m. – 10:30 a.m.	Part A – Overview of the Regulations
	Introductions

- Reasons for Change
- Structure of the Regulations
- Notified Bodies
- Transition Timelines
- Understanding Standards and Harmonization
- EU version of ISO 13485:2016 and ISO 14971:2007

Part B – MDR Article 120 Transition

- Status of the EU-MDR Implementation
- Article 120 Transitional Provisions
- Article 120
- ISO 13485:2016 & CEN/TR 17223:2018
- ISO 14971:2019
- Status of NBs
- Harmonized Standards & Common Specifications
- Status of Eudamed
- Manufacturer's Potential Hurdles
- Questions

Part C – Obligations of the Manufacturer

- Economic Operators
- Single Registration Number

	 QMS, RMS, and PMS CE Mark and Declaration of Conformity UDI requirements Incidents and Field Safety Corrective Actions Person Responsible for Regulatory Compliance Product Liability Insurance Exercise C1 – Person Responsible for Regulatory Compliance
	Exercise C2 – Economic Operator
10:30 a.m. – 10:45 a.m.	Break
10:45 a.m. – 12:00 p.m.	Part D – The Quality Management System (QMS)
	 ISO 13485:2016 & CEN/TR 17223:2018 QMS Requirements QMS Certificates Exercise D1 – Elements of the QMS
12:00 p.m. – 1:00 p.m.	Lunch Break
1:00 p.m. – 2:15 p.m.	Part E – The Risk Management System (RMS)
	 Requirements Description of Changes for ISO 14971:2019 The Process Flow in ISO 14971:2019 Annex I(3) The Risk Management System Annex I(4) Risk Control Measures Annex I(5) Use Error
2:15 p.m. – 2:30 p.m.	Break
2:30 p.m. – 5:00 p.m.	Part F – The Post-market Surveillance System (PMSS)
	 Risk Management Benefit-Risk Determination (BRD) Clinical Evaluation (CE) Post-market Clinical Follow-up (PMCF) Post-Market Surveillance Post-market Surveillance Report (PMSR) Periodic Safety Update Report (PSUR)

5:00 p.m.	 Summary of Safety and Clinical Performance (SSCP) Incident Reporting Exercise F1 – Elements of the PMSS And much more!
Day 3	
8:00 a.m. – 8:30 a.m.	Registration/Continental Breakfast
8:30 a.m. – 9:30 a.m.	Part A – The Internal Quality Audit Program
	 ISO 13485:2016 audit requirements Planning internal audits Conducting internal audits Reporting internal audits Resolving audit nonconformances
9:30 a.m. – 10:15 a.m.	Part B – EU-MDR/IVDR Conformity Assessment Paths
	 Annex VIII classification Options for EU-MDR Devices Options for EU-IVDR Devices Exercise B1 – Selecting a Conformity Assessment Path
10:15 a.m. – 10:30 a.m.	Morning Break
10:30 a.m. – 11:15 a.m.	Part C – EU-MDR QMS & RMS Considerations
	 Understanding Article 10 Understanding CEN/TR 17223:2018 Understanding ISO 13485:2016 Understanding ISO 14971:2019 Developing the Annex II and Annex III documentation Exercise C1 – Identify and Resolve Gaps from ISO 13485:2016
11:15 a.m. – 12:00 p.m.	Part D – Preparing the Application to the NB
	EU-MDR Applications

• EU-IVDR Applications

12:00 p.m. – 1:00 p.m. Lunch Break 1:00 p.m. – 2:15 p.m. Part E – The NB Approach The NB QMS requirements • Specific NB audit areas • Specific NB competence requirements • • EU-MDR Article 106 Expert Panels EU-IVDR Article 100 Reference Laboratories • Exercise E1 – Review Specific NB Audit Areas and Develop • Plans 2:15 p.m. – 2:30 p.m. **Afternoon Break** 2:30 p.m. – 4:00 p.m. Part F – Mapping the NB Approach to the Audit Program Specific audits • • Specific internal auditor competence Resolving audit nonconformances • Management review • Exercise F1 – Map the NB Audit to the Internal Audit Program • Exercise F2 – Report to Management Review • 4:00 p.m. **Workshop Completed**