



Webinar on Conformity Assessment for Medical Devices SaMD

Date: 29 August 2022

Time: 9:00 – 6:00 CT (10:00-7:00 EDT)

Hybrid event – Zoom + in person

Objectives. Provide an overview and practical implementation of IMDRF’s principles of conformity assessment for medical devices as well as a case study for software as a medical device.

TIME	AGENDA
9:00 – 9:05	Opening: Housekeeping Message Sandra Ligia González, Technical Secretariat, IACRC
9:05 – 9:10	Welcome Message Vesa Vuniqi, International Relations Specialist, US FDA Latin America Office
<i>CONFORMITY ASSESSMENT FOR MEDICAL DEVICES OVERVIEW AND PRACTICAL IMPLEMENTATION ROUNDTABLE</i>	
9:10 – 10:35	<p>Conformity Assessment for Medical Devices Moderator: Sandra Ligia González, IACRC</p> <p>Principles of Conformity Assessment for Medical Devices (GHTF/SG1/N78:2012)</p> <ul style="list-style-type: none"> - Erin Cutts, US FDA (10 min) <p>Practical Implementation of Conformity Assessment of Medical Devices - Panel</p> <ul style="list-style-type: none"> - Erin Cutts, US FDA (30 min) - Francisco Iran Cartaxo, ANVISA (30 min) <p>Q&A - 15min</p>
10:35-10:50	BREAK
10:50-12:05	<p>Practical Implementation of Conformity Assessment of Medical Devices – Panel (<i>continued</i>)</p> <p>Moderator: Patricia Pineda, US FDA</p> <ul style="list-style-type: none"> - Mukoil Romanos, INVIMA (20 min) - Carolina Magnatti, ANMAT (20 min) - Brenda Guadalupe Olvera, COFEPRIS (20 min) <p>Q&A – 15 min</p>

<p>12:05-12:50</p>	<p>Approach to Conformity Assessment for Medical Devices – The Industry Perspective</p> <p>Moderator: Sandra Ligia González, IACRC</p> <ul style="list-style-type: none"> - Fatemeh Razjouyan, Director of Regulatory Policy, International and Harmonization Global Regulatory Policy, Medtronic (15 min) - Duglas Rodríguez-Calderón, Head of LATAM Regulatory Policy, Global Regulatory Policy & Intelligence, Roche Diagnostics (15 min) <p>Q&A - 15min</p>
<p>12:50-2:00</p>	<p>LUNCH BREAK</p>
<p style="text-align: center;">CONFORMITY ASSESSMENT FOR MEDICAL DEVICES CASE STUDY</p>	
<p>2:00-3:05</p>	<p>Moderator: Patricia Pineda, USFDA</p> <p>Software as a Medical Device (SaMD) Key Definitions (IMDRF/SaMD WG/N12)</p> <ul style="list-style-type: none"> - Cathy Bar, US FDA (15 min) <p>Software as a Medical Device: Possible Framework for Risk Categorization and Corresponding Considerations (IMDRF/SaMD WG/N12)</p> <ul style="list-style-type: none"> - Brendan O’Leary, US FDA (15 min) <p>Software as a Medical Device (SaMD): Application of Quality Management System (IMDRF/SaMD WG/N23)</p> <ul style="list-style-type: none"> - Francisco Iran Cartaxo, ANVISA (15 min) <p>Q&A – 20min</p>
<p>3:05-4:25</p>	<p>Moderator: Sandra Ligia González, IACRC</p> <p>Practical Implementation of Conformity Assessment for SaMD</p> <ul style="list-style-type: none"> - MiRa Jacobs, US FDA (30 min) - Francisco Iran Cartaxo, ANVISA (30 min) <p>Q&A – 20 min</p>



4:25-4:40	BREAK
4:40-5:20	Approach to Conformity Assessment for SaMD – The Industry Perspective Moderator: Patricia Pineda, USFDA <ul style="list-style-type: none">- Diane Johnson, Sr. Director North American Policy, Global Digital Health Policy Lead, Johnson & Johnson MedTech (15min)- Duglas Rodríguez-Calderón, Head of LATAM Regulatory Policy, Global Regulatory Policy & Intelligence, Roche Diagnostics (15 min) Q&A – 10 min
5:20 – 5:30	Closing Remarks Vesa Vuniqui , US FDA/ Sandra Ligia González , IACRC