



2-day In-person Seminar:

Implementing ISO 13485:2016

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Boston, MA

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September 15th & 16th 2016

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9:00 AM to 6:00 PM



Dan O'Leary

President, Ombu Enterprises

Dan O'Leary is the President of Ombu Enterprises,

LLC, a company offering training and execution in Operational Excellence, focused on analytic skills and a systems approach to operations management. Dan has more than 30 years experience in quality, operations, and program management in regulated industries including aviation, defense, medical devices, and clinical labs. He has a Masters Degree in Mathematics; is an ASQ certified Biomedical Auditor, Quality Auditor, Quality Engineer, Reliability Engineer, and Six Sigma Black Belt; and is certified by APICS in Resource Management.

Overview:

The final version of ISO 13485:2016 is now available and companies should be planning their implementation. There are significant changes from the earlier version; they will require major modifications to the Quality Management System (QMS). The new version has better alignment with FDA's QSR, but there are still major differences that create issues for effective QMS implementation.

This workshop covers the differences from the 2003 version to the 2016 version and offers practical implementation advice to update your QMS. Participants will also learn the differences with QSR and understand how to resolve them. The workshop uses extensive examples and exercises to help clarify the concerns.

Some regulatory systems will rely on ISO 13485:2016 through the Medical Device Single Audit Program (MDSAP).

Price

(Without Stay) Price: \$1,295.00

(Seminar for One Delegate)

-(With Stay) Price: **\$1,695.00**

(Seminar for One Delegate)

Register now and save \$200. (Early Bird)

Register for 5 attendees (With stay)

Price: **\$4,323.00** You Save: \$4,152.0 (49%)*

\$8,475.00

ENROLL

**Please note the registration will be closed 2 days (48 Hours) prior to the date of the seminar.

Seminar Pricing Includes (With Stay)



Samsung Galaxy Tab 4



2 Days Stay



Pick-up and Drop Facility (Nearest Airport)



Break-Fast and Lunch



High Tea



Pack of 3 Webinars will be provided which has been done in the past on similar subject



Agenda:

Day One

Lecture 1: Overview of ISO 13485:2016

- Summary of the differences between ISO 13485:2003 and ISO 13485:2016
- · High level comparison with FDA QSR

Lecture 2: Regulatory Framework

- · Transition period for certificates
- · Canada MDSAP v CMDCAS
- EN ISO 13485:2016 and the MDD
- The new QMS audit findings/nonconformance grading system
- Implications of the EU's Medical Device Regulations

Lecture 3: Management Responsibility

- · Quality Policy and Objectives
- · Responsibility and Authority
- · Internal Quality Audits
- · Management Review

Lecture 4: Resource Management

- · Competence and Training
- · Infrastructure
- · Work Environment and Contamination Control

Lecture 5: Design and Development

- · Design Planning
- · Design Inputs and Design Outputs
- · Design Verification and Design Validation
- Design Review
- · Design Transfer
- Design Changes
- Design Files

Day Two

Lecture 1 : Supplier Management

- Selection
- · Purchasing Information
- · Written Quality Agreements
- · Purchased Product Verification

Lecture 2: Production Processes

- Production Control
- · Installation and Servicing
- · Identification and Traceability
- · Control of Nonconforming Product

Lecture 3: Process Validation

- · When to Validate
- · Validation Requirements
- Software Validation

Lecture 4: Monitoring and Measuring

- · Control of Equipment
- · Processes and Products
- Data Analysis
- · Complaint Handling

Lecture 5: Corrective and Preventive Action

- · Corrective Action
- Preventive Action



	Group Participation
10%	2 Attendees to get offer
20%	3 to 6 Attendees to get offer
25%	7 to 10 Attendees to get offer
30%	10+ Attendees to get offer

Payment Option

- 1 Credit Card: Use the Link to make Payment by Visa/Master/American Express card click on the register now link
- Check: Kindly make the check payable to NetZealous DBA GlobalCompliancePanel and mailed to 161 Mission Falls Lane, Suite 216, Fremont, CA 94539, USA
- 3 PO: Please drop an email to support@globalcompliancepanel.com or call the our toll free +1-800-447-9407 for the invoice and you may fax the PO to 302 288 6884
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What You will get

- 1 Learning Objectives
- 2 Participation certificates
- 3 Interactive sessions with the US expert
- 4 Post event email assistance to your queries.
- 5 Special price on future purchase of web based trainings.
- 6 Special price on future consulting or expertise services.
- 7 Special price on future seminars by GlobalCompliancePanel.
- 8 Seminar Kit includes presentation handout, ID card, brochure, trainings catalog, notepad and pen.
- 9 Networking with industry's top notch professionals

Contact Information: Event Coordinator

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Kindly get in touch with us for any help or information.

Look forward to meeting you at the seminar

GlobalCompliancePanel