




2-day In-person Seminar:

Quality by Design – Essential Techniques for Medical Devices

-  Salt Lake City, UT
-  April 13th & 14th, 2017
-  9:00 AM to 6:00 PM



Susanne Manz

Quality and Compliance Expert

Susanne Manz, MBA, MBB, RAC, CQA is an accomplished leader in the medical device industry with emphasis on quality, compliance, and Six Sigma. She has an extensive background in quality and compliance for medical devices from new product development, to operations, to post-market activities. While at industry leaders like GE, J&J, and Medtronic, Susanne worked in various world-wide roles including Executive Business Consultant, Worldwide Director of Quality Engineering, Worldwide Director of Design Quality, and Director of Corporate Compliance. She has traveled extensively throughout the world conducting audits and helping companies to understand and improve their Quality Management Systems.

Overview :

Design Controls are essential for producing safe and effective medical devices. And Design Controls are considered a critical process by the FDA. Yet, Design Controls are still one of the most frequent areas for 483 and Warning Letter observations. This 2 day seminar will help you understand, develop, and implement design controls processes and tools that are a competitive strength for your company. You will learn how to incorporate design controls into your product development process to help streamline development and ensure quality and compliance.

Price

Price: **\$1,495.00**

(Seminar for One Delegate)

Register for 5 attendees

Price: **\$4,485.00** You Save: \$2,990.0 (40%)*

~~\$7,475.00~~

ENROLL

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



Agenda:

Day One

Lecture 1:

- Overview and Expectations
- Background of FDA regulations
- Design Controls as an integrated part of New Product Development

Lecture 2:

- Design Planning
- Project Management
- Design Inputs

Lecture 3:

- Design Outputs
- Tools, forms, documents

Lecture 4:

- Design Verification and Validation
 - Concepts
 - Strategies
 - Statistical techniques

Why should you attend:

The intrinsic quality, safety, and effectiveness of medical device are established during the design phase. Yet, statistics show that a significant percentage of all medical device recalls are due to design problems. And those design problems can have disastrous results for your customer and for your company. A rigorous and efficient design control process can help avoid these quality and compliance problems. Issues that are identified early are more easily and quickly resolved.

Day Two

Lecture 1:

- Design Review
- Design History File
- Documentation Requirements

Lecture 2:

- Design Transfer
- Design for manufacturability concepts
- Design Changes
- Change Control and configuration management

Lecture 3:

- Linkages to other quality sub-systems
 - Risk Management
 - Failure Investigation
 - CAPA
- Inspection Preparedness and Compliance Strategy

Lecture 4:

- Lessons Learned
- Myths
- Challenges
- Best Practices

Who will benefit:

- R&D Engineers
- R&D Project Leaders
- R&D Managers and Directors
- Individuals participating in Product Design and Development
- Individuals participating in design changes and failure investigations
- Regulatory Affairs
- Design Quality Engineers
- R&D engineers and scientists
- Compliance Specialists

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
- 2** 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
- 4** Wire Transfer: Please drop an email to support@globalcompliancepanel.com or call our toll free +1-800-447-9407 for the wire transfer information

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

NetZealous LLC, DBA GlobalCompliancePanel

161 Mission Falls Lane, Suite 216,

Fremont, CA 94539, USA

Toll free: +1-800-447-9407

Fax: 302 288 6884

Email: support@globalcompliancepanel.com

www.globalcompliancepanel.com

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

Look forward to meeting you at the seminar

GlobalCompliancePanel