

Webinar on

Usability Engineering Under IEC 62366

Date : July 19, 2021

Areas Covered

The main focus will be on presenting tips and techniques for the practical application of the standard. Each phase in the development lifecycle will be discussed along with how deliverables required can be created, maintained, and evolved to meet regulatory requirements.



This training session will discuss techniques and methods that have proven successful in better ensuring safe and effective products and compliant regulatory submissions.

PRESENTED BY:

Don Hurd - has over 35 years of experience in supporting the development of applications of or containing software in regulated industries, the last 17 in medical devices. With his diverse background, Mr. Hurd provides a unique insight into driving product quality and ensuring high productivity of development organizations.



Date : July 19, 2021 Time : 01 : 00 PM EST Duration : 60 Minutes Price: \$179

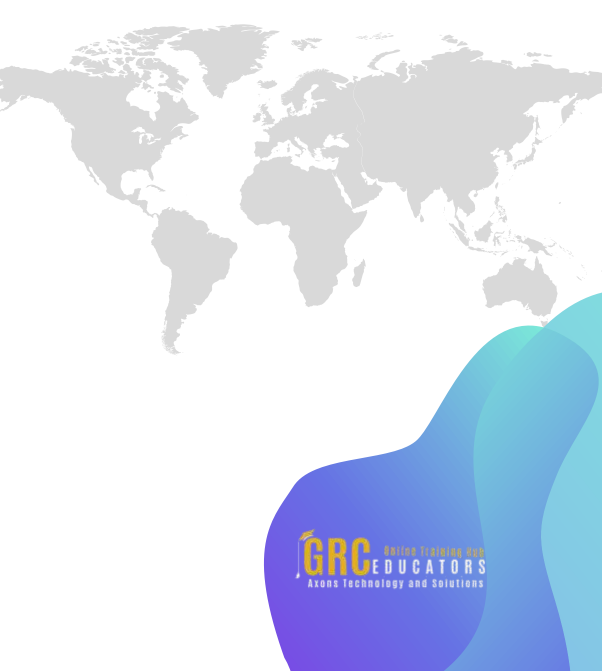
Webinar Description

ANSI/AAMI/IEC 62366 is a Recognized Consensus Standard by the US FDA and a harmonized standard in the EU. This means that compliance to it provides a presumption of conformity to the requirements within those jurisdictions. Compliance with the standard helps both ensure a smooth path to market clearance and drives a safer product.



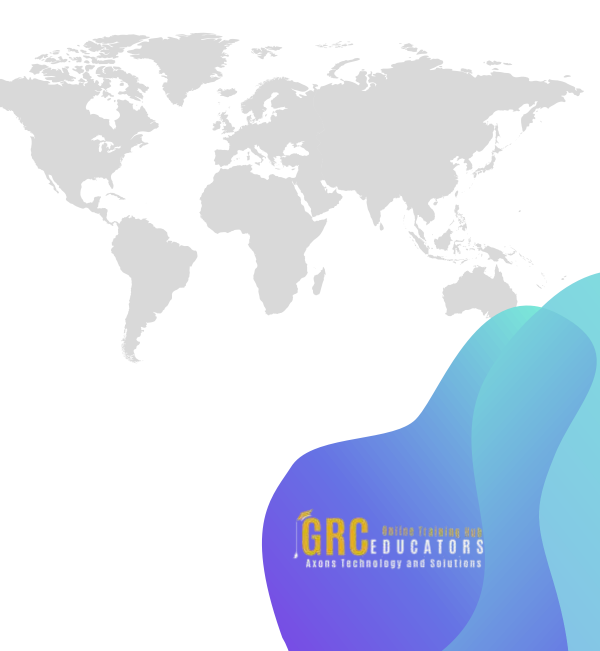
Who Should Attend ?

This webinar is intended for usability/human factors engineers, quality personnel, and project managers who manage projects with a user interface



Why Should You Attend ?

A practical application of the standard is not always intuitive. There are some nuances that can lead to unnecessary and excessive work. This training session will discuss techniques and methods that have proven successful in better ensuring safe and effective products and compliant regulatory submissions.



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