How to achieve compliance with IEC 62304 for medical device software development
- IEC 62304 Overview
- Standards Landscape
- Key elements of IEC 62304
- IBM Rational solution
- Recommendation
- Conclusion
IEC 62304 Overview

- **IEC 62304:2006 Medical device software – Software life cycle processes**

- Software use should not cause any unacceptable risk with respect to safety and effectiveness of the device

- Focused on software development and **maintenance** processes for medical devices but does **not** specify the methodologies, artifacts or life cycle models themselves

- Derived from ISO/IEC 12207, a general standard for software processes

- **Adoption**
  - FDA Consensus Standard since September 2008
  - FDA regards complying with IEC62304 as fulfilling “Software Development Environment Description” section of the *Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices*
  - Normative standard in Europe for conformance marking

- Standard available for purchase from ISO website (~$225 USD)
IEC 62304 Structure

- **Processes**: Set of interrelated or interacting activities that transforms inputs into outputs
- **Activities**: Set of interrelated or interacting tasks
- **Tasks**: Single piece of work that needs to be done and results in a deliverable
What IEC 62304 does not do

Does not cover validation and final release of a medical device

- Does not specify an organizational structure
  - You can do have a hierarchical, matrix, or mixed organization

- Does not specify the content of the documentation to be developed
  - You need to show traceability through all the artifacts but not in some set format

- Does not prescribe a specific lifecycle model
  - Waterfall, Iterative, Evolutionary, … it is all up to you
Standards Landscape

- Quality management system
- RISK MANAGEMENT
  - Software safety classification
- Software development PROCESS
  - Software development planning
  - Software requirements analysis
  - Software ARCHITECTURAL design
  - Software detailed design
  - SOFTWARE UNIT implementation and verification
  - Software integration and integration testing
  - SOFTWARE SYSTEM testing
  - Software release
- Software maintenance PROCESS
- Software RISK MANAGEMENT PROCESS
- Software configuration management PROCESS
- Software problem resolution PROCESS

Note: ISO 14971 is a Normative standard for Risk Management Process

Source: European Medical Device & Technology, June 2010
# IEC 62304 Software Development Lifecycle - Model

<table>
<thead>
<tr>
<th>DEFINITION / DEVELOPMENT</th>
<th>TEST / VERIFICATION</th>
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<tr>
<td><strong>System &amp; Stakeholder Requirements Definition</strong></td>
<td><strong>System Verification Test</strong></td>
</tr>
<tr>
<td>Output: System Requirements Doc. (SRD)</td>
<td>Output: System Verification Test Procedure (SVTP) and Results (SVTR)</td>
</tr>
<tr>
<td><strong>System Design &amp; Analysis</strong></td>
<td><strong>H/W - S/W Integration Test</strong></td>
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<tr>
<td><strong>Safety Classification</strong></td>
<td>Output: S/W Verification Cases and Procedures (SVCP) and Results (SVR)</td>
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<tr>
<td><strong>Output: Software Requirements Spec. (SRS)</strong></td>
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<td><strong>Output: Software Design Description (SDD)</strong></td>
<td><strong>Output: S/W Verification Cases and Procedures</strong></td>
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<tr>
<td><strong>Coding</strong></td>
<td><strong>Output: S/W Verification Cases and Procedures</strong></td>
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<tr>
<td><strong>Output: Source and Object Code</strong></td>
<td><strong>Output: S/W Verification Cases and Procedures</strong></td>
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<tr>
<td><strong>Code Review</strong></td>
<td><strong>Output: S/W Verification Cases and Procedures</strong></td>
</tr>
<tr>
<td><strong>Output: Source Code Review and Analysis Data</strong></td>
<td><strong>Output: S/W Verification Cases and Procedures</strong></td>
</tr>
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## Traceability

- **Traceability SRS to SRD**
- **Traceability SDD to SRS**
- **Traceability for Test Coverage**

## Configuration Management
- **Configuration Management**
- **Change Management (Problem Reporting)**
- **Quality Assurance Phase**
Software risk management in addition to ISO 14971 (see also IEC 80002 for specifics)

- Adds RISK CONTROL measures to Requirements
- Includes software to control MEDICAL DEVICE RISKS
- Allows for Design Risk Management (Bottoms Up)
- Complements Functional Risk Management (Top Down)
- Risk Management applied recursively throughout product lifecycle
- Documentation – various tools, safety assurance cases
Anticipate possible failures of the system
- Define control measures
  - Inherently safe
  - Preventive
  - Corrective
  - Informative

Systematic risk analysis is to anticipate failures
  - Top-down: Function analysis - ISO 14971, FTA
    - Hazard Analysis
  - Bottom-up: Design/ Process Analysis – FMEA
    - Failure Modes and Effects Analysis

Each failure leads to risk control (RCM) measures
Each RCM leads to requirements implemented in product hardware, software or documentation
Risk Management File documents traceably risk to control measure, to verification of control measure
Risk Management Activities continue after release
Document TRACEABILITY of software HAZARDS
From hazardous situation to the SOFTWARE ITEM
From SOFTWARE ITEM to the specific software cause
From the software cause to RISK CONTROL measure
From RISK CONTROL measure to VERIFICATION of RISK CONTROL measure

IEC 62304 Risk Management – Traceability, Verification
62304 Safety Classifications

“The software safety classes shall initially be assigned based on severity as follows:

- **Class A:** No injury or damage to health is *possible*
- **Class B:** Non-SERIOUS INJURY is possible
- **Class C:** Death or SERIOUS INJURY is possible”

Class C – 100% of 62304 activities apply
Class B – 93%
Class A – 43%
Software Systems are comprised of Software Items

Software Systems are assigned a Safety Classification

Software Items inherit the Safety Classification of the System

“If the HAZARD could arise from the failure of the SOFTWARE SYSTEM to behave as specified, the probability of such failure shall be assumed to be 100%”

Higher classification applies unless there is a documented rationale in a Risk Management file to justify lower safety class for software items
Segregation of Software Items
Safety, risk, and risk management process

- Safety is freedom from unacceptable risk.
- Safety is not security!
  - Security is protection of information and data so that unauthorized people or systems cannot read or modify them, and so that authorized people or systems are not denied access to them.

- Risk is a product of severity and probability of occurrence
- Software Risk Management Process:
  - identify software items that could contribute to a hazardous situation
  - identify causes of contribution to a hazardous situation
  - define risk control measures
  - verify risk control measures
  - Document traceability:
    - Hazardous situation to software item to specific software cause to risk control measure to verification of risk control measure
Hazard and Fault Tree Analysis

- Fault Tree Analysis determines what combinations of conditions or events are necessary for a hazard condition to occur

- Fault Tree Analysis allows the developer to identify, control and lower the risk to an acceptable level
  - This is critical in the IEC 62304 standard
Example Fault Tree Analysis
The key to safe systems is to analyze the system and to identify the conditions and events that can lead to hazards. Fault Tree Analysis (FTA) determines what logical combination of events and conditions lead to faults. By adding "ANDing-redundancy", architectural redundancy can be added.
Connecting FTA to Requirements (TraceToReq)
Manage Requirements across the Product Lifecycle
Capture, define, analyze, and manage requirements

- Improves visibility and collaboration
  - Accurate documentation for regulatory audits and reviews
- Supports FDA CFR21 Part 11 compliant electronic signatures
- Integrates with:
  - Portfolio management
  - Modeling
  - change management
  - Quality management solutions
Develop Systems and Software in a Model-Driven way

Visually develop complex systems using a structured approach

- Traceability from requirements -> implementation
- **Automate FDA documentation** submission
- Visual modeling manages complexity
- Generates production quality code
- Reduce testing time with model-driven testing
- Leverage existing code for documentation

Show link to IEC sections
Manage Change for Good Manufacturing Practice (GMP)
Establish an integrated change process across the lifecycle

- Manage Portfolio & Product Priorities
- Capture & manage requirements
- Develop Model-Driven System -> Software
- Capture customer requests & market driven enhancements
- Collaborate across Development Disciplines
- Integrate Suppliers
- Execute Tests

Collaboration, Process, Workflow

Integrated Change Management

Configuration Management

- Software
- Electrical
- Mechanical

Testing Eco-system

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Automate Document Generation
*Generate the right document at the right time*

- Increase productivity by allowing engineers to focus on engineering
  - NOT formatting concerns

- Maintain accuracy through quick one-click document generation
  - Captures last minute changes from data held in different source applications

- Enhance documentation quality thru template reuse
Recommendations

- Determine compliance with IEC 62304 by performing gap analysis
  - Create a chart that maps IEC 62304 clauses and subclauses to company procedure(s)
  - The mapping chart can be a standalone document, or it can be an appendix to a company’s procedure(s).

- Address open issues revealed in gap analysis
  - Establish missing processes
  - Amend deficient processes
  - Deploy appropriate software development tools that execute or enforce your processes

- Document updated software development process showing compliance with IEC 62304
  - Mapping chart should now have 100% coverage of IEC 62304 clauses within your processes
IBM Rational Software Platform for Systems
*Full coverage of the systems engineering lifecycle*

- Spans the entire systems and software lifecycle
- Integrates complex systems, embedded software and IT to create innovative products
- Achieves end-to-end traceability
- Provides proof of compliance
Waters Corporation
Testing equipment for medical, pharmaceutical and food products

What’s smart?

- Efficient systems for verifying the purity of drugs, food products and water resources
- Highly accurate blood tests with greater precision for healthcare diagnosis

Smarter business outcomes

- Innovation to enable significant advancements in healthcare delivery, environmental management, food safety, and water quality
- Increased quality and throughput of the assays performed with cost effective technology

How Rational enables smarter products

- Full traceability with an integrated requirements, change, and configuration management solution
- Performance improvement through global collaboration and component-based development

“After about 15 minutes of spending with the auditor, he was just blown away on how effective the Rational tools were in terms of addressing all of his audit questions.”
Customer Success: Safe Ventilation – at rest and on the move

*Hamilton Medical AG and IBM partner EVOCEAN GmbH*

What’s smart?
- Intelligent ventilation for intensive care
- Innovative cockpit display, ease of use
- Frees medical staff, improves patient outcome, reduces cost

Smarter business outcomes:
- Earlier error recognition, using modeling approach with code generation
- Consistent documentation with direct association between design and code
- Improved collaboration

How Rational Rhapsody helps:
- Proven embedded and real-time track record
- Code generation
- Re-usable Software Plattform

"Thanks to graphical representation of processes and states and being able to execute them, we now have a significantly improved basis for discussion. It's also easier for new employees to get going - provided they have some basic understanding of UML and Rhapsody®, “

Andreas Anderegg – Software Engineer
IBM Rational Solution for Medical Devices

**Execute best practices and collaborate through an integrated product lifecycle solution**

- Deliver medical devices that address market needs through portfolio management
- Provide clear audit trail across the development lifecycle with requirements management
- Validate designs early with model driven development
- Integrate change processes to coordinate development
- Automate document generation for compliance
- Drive quality throughout the product lifecycle

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TOP 5 REASONS TO ATTEND

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Innovate 2011 focuses on systems engineering and embedded software development on the "Modeling, Architecture and Construction track" and "new System Engineering track".

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Back-up
## Safety-Related Concepts

- **Accident** is a loss of some kind, such as injury, death, or equipment damage.

- **Risk** is a combination of the likelihood of an accident and its severity:
  \[
  \text{risk} = p(a) \times s(a)
  \]

- **A Hazard** is a set of conditions and/or events that leads to an accident.
  - Hazards are predictable and therefore controllable.
  - A safety-relevant system contains two kinds of hazards:
    - Intrinsic hazards - due to the inherent job and operational environment of the system.
    - Technology hazards - due to the addition of specific technological solutions.

- **A safety control measure** is an action or mechanism to improve the safety of the system by either:
  - Reducing the severity of the accident.
  - Reducing the likelihood of the accident.

- A fault is the nonperformance of a system or component and may be either random or systematic.

- Fault Tree Analysis determines what combinations of conditions or events are necessary for a hazard condition to occur:
  - Allows the developer to lower the risk of a Safety violation, critical for IEC 62304.
Safety Metamodel

- SafetyMeasure
  - faultActionTime: float
  - faultDetectionTime: float
  - faultTimeUnits: RhsString
  - safetyMechanism: RhsString

- UserModelClass
- FaultSource
  - faultMechanism: RhsString

- ManifestRelation

- Requirement
  - TraceToRequirement

- Hazard
  - faultToleranceTime: int
  - faultToleranceTimeUnits: RhsString
  - risk: float
  - severity: float
  - SafetyIntegrityLevel: RhsString

- ExterminatesRelation
  - exterminator

- FTANode
  - safetyConcern
  - probability: float

- LogicFlow
  - LogicFlow has exactly one association to a metatyp subtype of FTANode, either an output or an input.
  - UnaryOperator
  - LogicalOperator
  - BinaryOperator

- OutputNode
  - inputFlow
  - outputFlow

- Fault
  - MTBF: float
  - MTBF_TimeUnit: RhsString

- BasicFault
- UndevelopedFault

- ResultingCondition

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Fault Tree Analysis is defined in IEC 61025

Example of a Simplified Fault Tree Diagram for an Infusion Pump