IEC 62304 Software Development

A Real-World Perspective

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ISS Integrated Scientific Services AG
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# IEC 62304 Versions

<table>
<thead>
<tr>
<th>IEC 62304:2006 / EN 62304:2006</th>
<th>Edition 1.0</th>
<th>Harmonized for MDD</th>
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<td>IEC 62304:20xx</td>
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Basic Concepts

**Software** is anything that is executed on a processor or by an interpreter; it's the thing you cannot touch but which makes the other thing that you can touch do the thing you want*:

- Application Software on a PC
- Mobile App on a phone
- Firmware / Embedded Software inside a device
- FPGA Source Code – Configurable hardware that is “programmed” (as of Amd1:2015)

* the computer does what you tell it to do, not what you want it to do
Embedded software or Medical Device Software is software that is part of a physical medical device and is designed for a specific hardware.

Standalone software or Software as a Medical Device (SaMD) is software that is not part of a physical medical device and is designed to run on a generic hardware/software platform. The software is a medical device in itself. Where the platform is the combination of hardware and software that is not part of the health software, but is required to run it; e.g. operating system, system libraries.

Note: The Medical Device Coordination Group (MDCG) uses the term Medical Device Software (MDSW) for both, embedded and standalone software.
**SOUP (Software of Unknown Provenance)**

- Software item that is already developed and generally available and that has not been developed for the purpose of being incorporated into the medical device (also known as “off-the-shelf-software” or “3rd party component”).
- Software item previously developed for which adequate records of the development processes are not available

**Note:** Usually integrated into a product to reduce development time and cost and/or not to re-invent the wheel.

**Note:** The standard defines requirements and activities regarding the use of SOUP, adding SOUP thus also incurs some effort and costs.
Product Lifecycle

First loop is **development**
Subsequent loops are **maintenance**
Medical Software Lifecycle

Planning

Analysis

Requirements

Testing

Design

Implementation

Risk Management

Post Market

Decommission
Development Models

- Multiple ways to develop software exist, each one with advantages and disadvantages
- Not all are suited for all environments; e.g. medical devices
- People can get a bit religious about them
IEC 62304 / EN 62304 at a Glance

– The IEC 62304 is a process standard, it defines requirements to the development but not the product itself.

– Evidence of the correct application of the standard, i.e. performing the required activities, is the documentation

– Does not want to force a development model / process (e.g. Waterfall, V-model, SCRUM), but...
Different View (embedded software)
Different View (standalone software)
Coverage of IEC 62304 and IEC 82304-1
Risk Based Approach

The standard defines three software safety classes:

− **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

The software safety class applies to the software as a whole, but may optionally be applied to individual or groups of software modules or components, e.g. low-risk components may fall into a lower safety class.

The software safety class is defined by the potential harm emanating from the software, regardless of the probability of the harm actually occurring.

**Note:** Software safety class is independent from the medical device classification (Europe) and the level of concern (FDA); all of them are risk-based though.
The software safety class determines which lifecycle activities need to be performed for the specific part of the software:

<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Class A</th>
<th>Class B</th>
<th>Class C</th>
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<tr>
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<td>Software development PROCESS</td>
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<td>5.1.7</td>
<td>Software RISK MANAGEMENT planning</td>
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The software safety class should be determined through risk and hazard analysis:

<table>
<thead>
<tr>
<th>Probability</th>
<th>Residual risks only related to device software</th>
<th>Severity</th>
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<td></td>
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<td>Class A</td>
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Software Safety Classification

– As early as possible as it determines the activities required
– If not known (e.g. risk management not done yet) assume safety class C
– For the residual risk, only consider risk control measures external to the software
  The standard essentially assumes that if the software fails, it fails catastrophically, i.e. negating software-based risk control measures.

Note: The standard states that “Probability of a software failure shall be assumed to be 1”, this only concerns the software safety classification and not failure probability for risk management or in general
Software Safety Classification

Software items with a lower safety class must not be able to adversely affect software items of a higher safety class; the manufacturer must provide rationale for the segregation between such items. If this is not possible, the items cannot be in a lower safety class.

Also software-based risk control measures take the safety class of the risk they’re controlling.
Processes defined by the IEC 62304:

- Software development (section 5)
- Software maintenance (section 6)
- Software risk management (section 7)
- Software configuration management (section 8)
- Software problem resolution (section 9)
Software Development Process

Activities outside of the scope of this standard

Customer needs satisfied

Customer Needs

System development activities (including risk management)

7 Software risk management

5.1 Software development planning
5.2 Software requirements analysis
5.3 Software architectural design
5.4 Software detailed design
5.5 Software unit implementation and verification
5.6 Software integration and integration testing
5.7 Software system testing
5.8 Software release

8 Software configuration management

9 Software problem resolution
Software Maintenance Process

6.1 Establish software maintenance plan
6.2 Problem and modification analysis
5.3 Software architectural design
5.4 Software detailed design
5.5 Software unit implementation and verification
5.6 Software integration and integration testing
5.7 Software system testing
5.8 Software release
6.3 Modification implementation
7 Software risk management
8 Software configuration management
9 Software problem resolution

Activities outside of the scope of this standard
System maintenance activities (including risk management)
Request satisfied
Maintenance Request
Request satisfied
Software Risk Management Process

Risk Management with a software perspective:

- Identify software items that could contribute to a hazardous situation and identify the potential causes
- Define and implement risk control measures
- Verify risk control measures (implemented and effective)
- Performed during development and maintenance activities; e.g. design changes must be analyzed for their impact on the patient risk (new risks, impact on existing risks or risk control measures)

**Note:** IEC 62304 mandates risk management according to ISO 14971

**Note:** Software developers are not risk managers! They can and should be part of the risk management team as subject matter experts though.
**Configuration Management Process**

Identify the inputs (and environment) from which a specific output has been developed:

- Establish means to **identify configuration items** (traceability)
- Techfile documents as defined in the **document plan**
- Source code as maintained in the **version control system**
- Toolchain, SOUP, and other inputs as defined in the **configuration manual**

**Note:** The configuration of a software release is essentially a snapshot of all the inputs and the generated output, including all relevant documents of the techfile.
Problem Resolution Process

Handle problems found during development or in the field:

- Prepare **problem report** for each problem found
- Investigate the problem and analyze the problem’s **impact on safety** (risk management, not the developer)
- Inform relevant parties
- Create **change requests** (change management, maintenance process) to fix the problem
- Verify problem resolution (also add to regression tests)
- Analyse problems for trends

**Note:** Most of these activities can be handled by a properly configured ticketing system.
Chronological order often encountered in projects

- Configuration management is neglected during development and maintenance and rushed before a release (document signing marathon)
- Problem resolution processes usually start when the software release is almost done and fails some minor tests; e.g. defects are entered into the ticketing system and fixed in a future release
IEC 62304 in the Field, as seen by a Software Development Consultant

Types of Projects

− Green Field vs Brown Field
− Motivated vs non-Motivated
− Fast Exit vs becoming Legal Manufacturer
− Development processes applied vs “uh, something like scrum?”

Note: Usually people only call for external help when they’re already stuck
Example – “University Project”

- Develops a proof of concept and wants to market it
- High employee turnover (e.g. built by postdocs and research assistants)
- Does not employ a structured development process
- No experience in medical device development / medical device software development
- No reasonable way to make the existing result compliant with the regulatory or normative requirements
- Project abort or redo from start by an experienced supplier or manufacturer

Note: To make things worse, proof of concept sometimes already used on patients through university clinics or similar institution.
Example – “Startup, not motivated”

- Develops a product and wants to market it
- Suddenly confronted by fact that product is a medical device and covered by legislative and normative requirements (and that these also apply to startups)
- Goal is quick exit and not becoming a legal manufacturer
- Retrospective application of IEC 82304/IEC 62304 activities and documentation; find gaps and fill them
- IEC 82304/IEC 62304 outsourced / ghostwriting by service provider
- No or negligible internal acquisition of know-how
- Painful but possible for Class I products (MDD, likely no longer possible under MDR)
Retrospective documentation and execution of lifecycle activities

Activities outside of the scope of this standard

Customer needs satisfied

System development activities (including risk management)

7 Software risk management

5.5 Software unit implementation and verification

5.1 Documentation of the software development

5.2 Documentation of the software requirements

5.3 Documentation of the software architecture

5.4 Documentation of the detailed software design

5.6 Retrospective software integration and verification

5.7 Retrospective software system testing

5.8 Retrospective software release

8 Software configuration management

9 Software problem resolution

Solution looking for a problem (profitable Exit desired)

Customer Needs (if any)
Example – “Startup, motivated”

− Develops a product and wants to market it
− Suddenly confronted by fact that product is a medical device and covered by legislative and normative requirements
− Goal is a medium- or long-term exit (e.g. after clinical trial) or becoming a legal manufacturer
− Retrospective application of IEC 82304/IEC 62304 activities and documentation; find gaps and fill them
− Introduction of quality management system
− Requires external help, often also to acquire the required knowhow
− May require changes to the product
− Painful but possible
Example – “Legacy”

– Existing product, possibly on the market for a long time
– New features or regulatory changes makes it a medical device
– Retrospective application of IEC 82304/IEC 62304 activities and documentation; find gaps and fill them; reverse engineering may be required (legacy process)
– Introduction of quality management system or transformation / extension of existing QMS (e.g. ISO 9001 to ISO 13485)
– Usually requires external help, often to acquire the required knowhow
– Painful but possible

Note: The legacy software process can only be applied for software that has already been placed on the market legally.
What does work

– Compliant development from the start
– Retrospective activities and documentation for legacy projects or motivated startups
– Remove medical functionality and market product as lifestyle product or medical device of a lower class
– Using the time the product is sold as a lifestyle product to get the required processes and infrastructure to create medical devices, collecting and using post market data from the lifestyle product and in the end add the medical device functionality back to the product and release it as such
– Moving from pure agile development to IEC compliant development
What usually doesn’t work

- Shadow-Techfile / Ghostwriting
- University → CE Mark / University as a legal manufacturer
- Pure agile software development (e.g. pure SCRUM)
- Forcing compliant software development processes on developers that don’t want to use them and/or don’t understand the reasons (e.g. when moving from SCRUM to IEC 62304 development processes)
- Replacing consultants until you find one that tells you that you don’t have to do change anything…
Medical Device Software Development – The Right Mindset

– The goal is to achieve the intended use with minimal risk to the patient
– Processes are here for a reason and not just to torture the developer
– Software development is one part of a whole; no lone guns; changes to the software design may have impact on e.g. risk management, usability engineering, etc…
– Design changes are not local; no tunnel vision
– Error handling vs Essential Performance
– Design changes only through the software maintenance process
– SOUP is not free (there ain’t no such thing as a free lunch)
– SOUP – don’t forget software licenses (e.g. open source)
– Technical debt is even worse than for “normal software”
IEC 62304 and Agile Development

While the IEC 62304 does not want to promote a specific development model, it strongly leans towards the V-model. Today software is often developed iteratively (agile software development); which can be done compliant to the IEC 62304:

- Interpret the V-model of the standard as document landscape and not as fixed development process
- Create and release software development plan and document plan at the start of the development
  - Update draft versions continuously
  - Release as new version on major changes
- Continuously update all documents, software requirements and design must be released before testing (Verification)
- Plan, perform and document reviews
- Integrate risk management into processes
IEC 62304 and Agile Development

- The scope of the IEC 62304 is wider than that of most agile methodologies. The agile process need to be expanded with the additional activities
- The AAMI Technical Report TIR 45 provides further information on how to implement IEC 62304 compliant agile development
- Human factor; don’t leave the developers behind
A quick look into the crystal ball
Cybersecurity – IEC CD 80001-5-1 Cyber Security Lifecycle (draft)

- Draft for IEC 80001-5-1 that would extend IEC 62304 software lifecycle processes with security related activities
- Current estimate for publishing is mid-2021
Cybersecurity – IEC CD 80001-5-1 Cyber Security Lifecycle (draft)

Customer Needs

Activities outside of the scope of this standard

System development activities (including risk management)

7 Software risk management

Security – Activities in the product lifecycle

5.1 Software development planning
5.2 Software requirements analysis
5.3 Software architectural design
5.4 Software detailed design
5.5 Software unit implementation and verification
5.6 Software integration and integration testing
5.7 Software system testing
5.8 Software release

Security – Activities in the product lifecycle

8 Software configuration management

Security – Activities in the product lifecycle

9 Software problem resolution

Security – Activities in the product lifecycle
IEC 62304 Edition 2

- Second draft has been rejected too
- Current estimate for publishing is mid-2021