Product Development for Medical, Life Sciences and Consumer Health
Usability Testing for FDA Validation

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Topics

- Why is usability testing required for FDA validation?
- How is validation testing different from other testing?
- Conducting the test (infusion pump example)
- Preparing the FDA report
- Case study: Drug delivery device
- Relevant standards and regulations
Why is usability testing required?
Why is usability testing required?

Testing for ease and accuracy of use is the **only way to ensure** that users can **safely and effectively** operate, install, and maintain devices.

This process **culminates in full testing** of a model embodying all the user-interface characteristics for both hardware and software of a fully functioning device.

-- from the FDA (1996), *Do It By Design*
Subpart C - Design Controls

The need for human factors techniques or data in the design process is implicit in paragraphs c, f, and g of Section 820.30.

(g) Design validation: "Design validation shall ensure that devices conform to defined user needs and intended uses, and shall include testing of production units under actual or simulated use conditions."

Human factors relevance: Design validation should be used to demonstrate that the potential for use error that can lead to patient injury has been minimized. The regulation requires testing the device under actual or simulated use conditions. Realistic use conditions, therefore, should be carried out by test participants who represent a range of typical intended users in terms of their ability to acquire information from, manipulate, and maintain the device and understand the accompanying labeling.

From the FDA -- “Human Factors Implications of the New GMP Rule Overall Requirements of the New Quality System Regulation”
This is the **most important** HFE report that the FDA reviews.

Iterative, formative usability testing is already done.

It should **tie back to**:

- User profiles
- FMEA
- Usability goals

It must validate the **safety and usability** of the device.
Conducting the validation test
Designing the test

Select & recruit participants
Select tasks or features to be tested
Select measures
Prepare the test materials
Handle logistics & preparation
Run a pilot test
What are some user profiles for a test of an infusion pump?
What are some **user profiles** for a test of an infusion pump?

- Nurses – specify
- Physicians – specify
- Others?
How many participants?

No specific number required by the FDA

**Sample size** should be larger than for formative tests

Consider the impact of potential **use errors**

Consider the **variability** of patient population

Ron Kaye of the FDA\(^1\):

“Upwards of 25 participants [per user group] seems about right...Rarely does small sample size arise as FDA’s primary reason for viewing a particular summative human factors test as unsatisfactory. Rather it is that the selected user tasks exclude some of the critical ones, the participants do not effectively represent the user population, the performance measurements are not germane, or there is no clear link between the test results and safety.”

Selecting tasks

**Primary operating functions** (POFs)

Subset - frequently performed functions

Address a **safety concern**

Performed under stress

**Redesigned** after previous testing

Illustrate error recovery
Example

What are some **tasks** you might test on an infusion pump?
Example

What are some tasks you might test on an infusion pump?

- Turn on the pump
- Set it up for drug administration by keying in the flow rate and volume-to-be-infused parameters
- Install the administration set
- Perform primary and secondary infusion
- Change parameters
- React to an emergency
- Set alarms
- Silence alarms
Selecting measures

**Objective** measures
- Measures of performance
- Observable

**Subjective** measures
- Ratings, opinions, comments

Base these on your *usability goals*
Which ones can be quantified?
Example

What are some measures you might use when testing an infusion pump?
Example

What are some measures you might use when testing an infusion pump?

- Task times
- Number of errors
- Types of errors
- Completed tasks (%)
- Usability ratings
- Failure to react to alarms
- Satisfaction with control layout
Conducting the sessions

**Where** to conduct the test?

How long should sessions take?

Who should **moderate**?

Should participants **think aloud**?
Conducting the sessions

**Where** to conduct the test?

How long should sessions take?

Who should **moderate**?

Should participants **think aloud**?

**Typical outline:**

- Introduction & informed consent
- Pre-test questionnaire
- User performs tasks
  - Objective & subjective data collected
- Post-test interview and ratings
Preparing the FDA report
Analyzing the data

Analyze **quantitative** data
- Calculate descriptive stats

Analyze **qualitative** data
- Moderator’s & observers’ notes
- Questionnaires & rating sheets
- Pre- and post-test interview results

Identify **usability issues**

Identify any residual **safety risks**
Examples of usability issues

Nurses *can’t figure out* how to insert tubing in an infusion pump

Lab technicians load disposables into the wrong drawer

Nurses *incorrectly program* parameters in a patient monitor

Patients misinterpret the LEDs on a ventricular assist device

Surgeons experience *pinch points* with a hand-held instrument
What constitutes a **successful task outcome**? It is difficult to establish a criterion for pass-fail rates 90%? 95%? 99%? Each error/failure must be carefully analyzed – does it place the user or patient at risk?

**Example:**

For this study, at least 28 of the 30 subjects must have registered a “success” for a particular scenario for the design to pass usability validation. The design must also score an average of 4.0 or higher on a 5-point scale in order to pass usability validation.
Report format

There is no prescribed format

**Tie it back** to the FMEA

Was usability **validated** or not?

Are additional design changes needed?
There is no prescribed format

Tie it back to the FMEA

Was usability validated or not?

Are additional design changes needed?

**Typical outline:**

- Purpose
- Methodology
- Data analysis
- Task success rates
- Errors and usability issues
- Conclusions (pass/fail)
Is re-testing needed?

Depends on what the **errors were** and their severity

If design changes were significant, you need to re-test

Often you can use a **smaller sample**

Sometimes you can re-test just the design changes
Case study: Drug delivery device

Client submitted for 510(k) clearance, but FDA required validation testing

**User profiles:** Patients and health care professionals

**Selected tasks:** All primary operating functions

**Measures:** Objective and subjective, based on hazards analysis

**Materials:** Two detailed moderator’s scripts, training, IFUs

**Test environment:** Market research facility

**Data collection:** Real-time using carefully designed spreadsheets

**Analysis:** Completed by client, reviewed by Farm

**Results:** Showed hazards had been addressed, no new issues
Final thoughts

Usability testing should be **iterative** during the design process.

Validation testing is **required** for CE Mark & FDA.

Validation testing must include **safety-critical** tasks.

Validation testing must be done with **actual users** in **actual or simulated** environments.
Standards & regulations

FDA Quality System Regulations
21 CFR 820.30, Subpart C
Design Controls, paragraphs c, f, and g

ANSI/AAMI HE74:2001
Human Factors Design Process for Medical Devices

ANSI/AAMI HE75:2009
Human Factors Engineering – Design of Medical Devices

ISO/IEC 62366:2008
Application of Usability Engineering to Medical Devices

Do It By Design (FDA, 1996)
An Introduction to Human Factors in Medical Devices
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THANK YOU.

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