

# Usability Engineering File

Use this template to iEC 62366 usability engineering file for [device\_name].

## Template Metadata

| Field          | Details                                      |
|----------------|--|
| Category       | Medical Devices                              |
| Owner          | [Team or owner]                              |
| Version        | [Version number]                             |
| Effective Date | [Date]                                       |
| Review Cycle   | [Monthly / Quarterly / Annual / Event-based] |
| Status         | [Draft / In Review / Approved]               |

## Use Specification

Define intended users, patient population, use environments, indications, contraindications, and operating principle.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

## Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## User Interface Characteristics

Describe controls, displays, alarms, packaging, labeling, software screens, and accessories.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

### Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Hazard-Related Use Scenarios

List critical tasks, foreseeable use errors, hazardous situations, harms, and risk controls.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

### Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Formative Evaluations

Summarize formative studies, participants, findings, and design changes.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

## Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Summative Validation

Describe validation method, participant groups, scenarios, success criteria, and observed use errors.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

## Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Residual Use Risk

Evaluate residual risks, labeling controls, benefit-risk rationale, and required mitigations.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

## Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Traceability

Map user needs, use scenarios, risk controls, and validation evidence. Use IEC 62366-style terminology and keep traceability explicit.

| Item                  | Details   | Owner   | Status            |
|-----------------------|---|---------|-------------------|
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |
| [Item or requirement] | [Describe the relevant detail, evidence, or decision] | [Owner] | [Open / Complete] |

## Notes

[Add context, assumptions, exceptions, evidence links, screenshots, calculations, or reviewer comments.]

## Review and Signoff

Document review conclusions, approvals, unresolved items, and next review date.

| Role     | Name   | Date   | Notes   |
|----------|--------|--------|---------|
| Preparer | [Name] | [Date] | [Notes] |
| Reviewer | [Name] | [Date] | [Notes] |
| Approver | [Name] | [Date] | [Notes] |