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## Cardiovascular implants and extracorporeal systems — Cardiac valve repair devices

*Implants cardiovasculaires et circuits extra-corporels — Dispositifs de réparation de valves cardiaques*

~~Second edition~~

~~Date: 2023-07-26~~

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### Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO document should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see [www.iso.org/directives](http://www.iso.org/directives)).

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For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT), see [www.iso.org/iso/foreword.html](http://www.iso.org/iso/foreword.html).

~~accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).~~

This document was prepared by ISO/TC 150, *Implants for surgery*, Subcommittee SC 2, *Cardiovascular implants and extracorporeal systems*.

This second edition cancels and replaces the first edition (ISO 5910:2018), which has been technically revised.

The main changes are as follows:

- ~~added normative~~ requirements (~~7.2.2.3~~) in 7.2.2.3 and ~~an~~ informative annex (~~Annex I~~) ~~on~~ have been added for test platforms;
- ~~removed normative~~ requirements and ~~the~~ informative annex on functional performance assessment have been removed;
- ~~combined~~ fatigue and durability testing have been combined into integrated device assessment (~~7.2.4~~ in 7.2.4 and ~~Annex M~~ Annex M);
- ~~moved~~ the description of ~~the~~ types of heart valve repair devices, including ~~the~~ added section on robotically-assisted systems, have been moved into ~~Annex B~~ Annex B;
- ~~Annex G~~ extensively revised annexes on hazard and failure modes (~~Annex G~~) and ~~Annex R~~ on clinical imaging (~~Annex R~~) have been extensively revised;
- ~~clarified~~ pulsatile flow conditions for the paediatric population have been clarified in ~~Annex H~~ Annex H;

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- ~~Annex I~~ ~~added annex covering~~ ~~on~~ boundary conditions ~~(Annex I)~~; ~~has been added~~;
- ~~removed~~ annexes on physical and material property definitions and material property testing ~~have been removed~~;
- ~~included~~ additional device design evaluation requirements ~~have been included in Annex N~~ ~~(Annex N)~~;
- ~~included~~ additional evaluation considerations for preclinical in vivo evaluations ~~have been included in Annex P~~ ~~(Annex P)~~.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at [www.iso.org/members.html](http://www.iso.org/members.html).

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# Cardiovascular implants and extracorporeal systems — Cardiac valve repair devices

## 1 Scope

~~1.1 This document is applicable to all heart valve repair systems that have an intended use to repair and/or improve the function of native human heart valves by acting either on the valve apparatus or on the adjacent anatomy (e.g. ventricle, coronary sinus).~~

~~1.2~~**1.1** This document specifies an approach for verifying and validating the design and manufacture of a heart valve repair system through risk management. The selection of appropriate verification and validation tests and methods are derived from the risk assessment. The tests include assessments of the physical, chemical, biological and mechanical properties of components and materials of heart valve repair systems. The tests also include preclinical in vivo evaluation and clinical investigation of the finished heart valve repair system to assess the safety and effectiveness of the heart valve repair system.

NOTE For the purposes of this document, effectiveness ~~endpoint~~ **end point** includes clinical performance and benefits.

~~1.3~~**1.2** This document defines operational conditions and performance requirements for heart valve repair systems where adequate scientific and/or clinical evidence exists for their justification. It also describes the labels and packaging of the device.

~~1.3 This document applies to all heart valve repair systems that have an intended use to repair and/or improve the function of native human heart valves by acting either on the valve apparatus or on the adjacent anatomy (e.g. ventricle, coronary sinus).~~

~~1.4~~**1.4** This document ~~is does not applicable~~ **apply** to cardiac resynchronization therapy (CRT) devices, paravalvular leakage closure devices, systems that do not leave an implant in place (e.g. ablation, radio frequency annuloplasty), apical conduits and devices with components containing viable cells. This document also excludes materials not intended for repairing and/or improving the function of human heart valves according to its intended use (e.g. patch material and sutures used in general surgical practice).

NOTE A rationale for the provisions of this document is given in [Annex A](#).

## 2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 10993-1, *Biological evaluation of medical devices — Part 1: Evaluation and testing within a risk management process*

ISO 10993-2, *Biological evaluation of medical devices — Part 2: Animal welfare requirements*

[ISO 10993-4, \*Biological evaluation of medical devices — Part 4: Selection of tests for interactions with blood\*](#)

ISO 11135, *Sterilization of health-care products — Ethylene oxide — Requirements for the development, validation and routine control of a sterilization process for medical devices*

ISO 11137-1, *Sterilization of health care products — Radiation — Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices*

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- ISO 11137-2, *Sterilization of health care products — Radiation — Part 2: Establishing the sterilization dose*
- ISO 11137-3, *Sterilization of health care products — Radiation — Part 3: Guidance on dosimetric aspects of development, validation and routine control*
- ISO 11607-1, *Packaging for terminally sterilized medical devices — Part 1: Requirements for materials, sterile barrier systems and packaging systems*
- ISO 11607-2, *Packaging for terminally sterilized medical devices — Part 2: Validation requirements for forming, sealing and assembly processes*
- ISO 13485, *Medical devices — Quality management systems — Requirements for regulatory purposes*
- ISO 14155, *Clinical investigation of medical devices for human subjects — Good clinical practice*
- ISO 14160, *Sterilization of health care products — Liquid chemical sterilizing agents for single-use medical devices utilizing animal tissues and their derivatives — Requirements for characterization, development, validation and routine control of a sterilization process for medical devices*
- ISO 14630, *Non-active surgical implants — General requirements*
- ISO 14937, *Sterilization of health care products — General requirements for characterization of a sterilizing agent and the development, validation and routine control of a sterilization process for medical devices*
- ISO 14971, *Medical devices — Application of risk management to medical devices*
- ISO 15223-1, *Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements*
- ISO 15223-2, *Medical devices — Symbols to be used with medical device labels, labelling, and information to be supplied — Part 2: Symbol development, selection and validation*
- ISO 17664-1, *Processing of health care products — Information to be provided by the medical device manufacturer for the processing of medical devices — Part 1: Critical and semi-critical medical devices*
- ISO 17665-1, *Sterilization of health care products — Moist heat — Part 1: Requirements for the development, validation and routine control of a sterilization process for medical devices*
- ISO/TS 17665-2, *Sterilization of health care products — Moist heat — Part 2: Guidance on the application of ISO 17665-1*
- ISO/TS 17665-3, *Sterilization of health care products — Moist heat — Part 3: Guidance on the designation of a medical device to a product family and processing category for steam sterilization*
- ISO 20417, *Medical devices — Information to be supplied by the manufacturer*
- ISO 22442-1, *Medical devices utilizing animal tissues and their derivatives — Part 1: Application of risk management*
- ISO 22442-2, *Medical devices utilizing animal tissues and their derivatives — Part 2: Controls on sourcing, collection and handling*
- ISO 22442-3, *Medical devices utilizing animal tissues and their derivatives — Part 3: Validation of the elimination and/or inactivation of viruses and transmissible spongiform encephalopathy (TSE) agents*

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ISO/TR 22442-4, *Medical devices utilizing animal tissues and their derivatives — Part 4: Principles for elimination and/or inactivation of transmissible spongiform encephalopathy (TSE) agents and validation assays for those processes*

IEC 62366-1, *Medical devices — Part 1: Application of usability engineering to medical devices*

[ASTM F1830, Standard Practice for Selection of Blood for In Vitro Evaluation of Blood Pumps](#)

### 3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminology databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <https://www.iso.org/obp>
- IEC Electropedia: available at <https://www.electropedia.org/>

#### 3.1

##### accessory

device-specific tool that is required to assist in the implantation and/or adjustment of the *heart valve repair device* (3.25), excluding the *delivery system* (3.12)

#### 3.2

##### active comparator

intervention (e.g. transcatheter or surgical valve repair or replacement) generally accepted as the standard of care for the intended valve device indication that can be used as a basis of comparison for the *safety* (3.45) and effectiveness of the *heart valve repair device* (3.25)

#### 3.3

##### adverse event

##### AE

untoward medical occurrence, unintended disease or injury, or untoward clinical signs (including abnormal laboratory findings) in subjects, users or other persons, whether or not related to the investigational medical device and whether anticipated or unanticipated

Note 1 to entry: This definition includes events related to the investigational medical device or the *active comparator* (3.2(3.2)).

Note 2 to entry: This definition includes events related to the procedures involved.

Note 3 to entry: For users or other persons, this definition is restricted to events related to the use of investigational medical devices or comparators.

#### 3.4

##### boundary ~~conditions~~condition

~~displacements, forces, moments, pressures, displacement, force, moment, pressure~~ or constraint ~~condition~~ acting on an implanted valve repair device and/or surrounding tissue

#### 3.5

##### cardiac output

##### CO

stroke volume [i.e. volume of blood pumped by a ventricle during systolic duration (3.53(3.51) times)] multiplied by the heart rate

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### 3.6

#### coating

thin-film material that is applied to an element of a *heart valve repair system* (3.26) to modify its surface physical or chemical properties

### 3.7

#### compliance

$C$  relationship between the change in radius and the change in pressure of a deformable tubular structure (e.g. valve annulus, aorta, conduit), defined given as

$$= \frac{(r_2 - r_1) \times 100}{r_1 \times (p_2 - p_1)} \times 100 \%$$

where

-  $C$  is the compliance in units of % radial change/100 mmHg;

$p_1$  is the diastolic pressure, in mmHg;

$p_2$  is the systolic pressure, in mmHg;

$r_1$  is the inner radius at  $p_1$ , in millimetresmm;

$r_2$  is the inner radius at  $p_2$ , in millimetresmm

Note 1 to entry: ~~Definition and equation adapted from ISO 25539-1~~ Compliance is expressed in percentage of radial change per 100 mmHg.

### 3.8

#### component-joining material

material, such as a suture, adhesive or welding compound, used to assemble the components of a *heart valve repair device* (3.25(3.25)), thereby becoming part of the implanted device

### 3.9

**cycle** <https://standards.iteh.ai/catalog/standards/iso/edc865d4-cb05-43d6-8413-b8f0ac5c4c36/iso-fdis-5910> complete sequence in the action of a native or repaired heart valve under pulsatile flow conditions

### 3.10

#### cycle rate

#### beat rate

number of complete *cycles* (3.9) per unit of time

Note 1 to entry: The cycle rate is usually expressed as cycles (3.9) per minute (cycles/min or beat/min); beats per minute.

### 3.11

#### delivery approach

anatomical access used to deliver the *implant* (3.28) to the *implant site* (3.29) (e.g. transfemoral, transapical, transeptal)

### 3.12

#### delivery system

catheter or other system used to deliver, deploy, attach or adjust the device in the *implant site* (3.29(3.29)).

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3.13

**design validation**

establishment by objective evidence that device specifications conform with user needs and *intended use(s)* [\(3.31\)](#) ~~(3.31)~~

3.14

**design verification**

establishment by objective evidence that the design output meets the design input requirements

3.15

**device ~~embolisation~~ embolization**

dislodgement from the intended and documented original position to an unintended and nontherapeutic location

3.16

**device failure**

inability of a device to perform its intended function

3.17

**device migration**

detectable movement or displacement of the *implant* [\(3.28\)](#) from its original position within the *implant site* [\(3.29\)](#), without device *embolization* [\(3.15\)](#) ~~*embolisation* [\(3.15\)](#)~~

3.18

**failure mode**

mechanism of *device failure* [\(3.16\)](#) ~~(3.16)~~

~~Note 1 to entry: EXAMPLE~~ Support structure [\(3.52\)](#) ~~fracture~~ [\(3.22\)](#), calcification, and prolapse ~~are examples of failure modes.~~

3.19

**fatigue**

process of progressive localized permanent structural change occurring in a material subjected to conditions that ~~produce~~ *produces* fluctuating stresses and strains at some point ~~or points(s)~~ and that can culminate in cracks or complete *fracture* [\(3.22\)](#) after a sufficient number of fluctuations

3.20

**follow-up**

continued assessment of subjects who have received the *heart valve repair device* [\(3.25\)](#)

3.21

**forward flow volume**

volume of flow ejected through the repaired heart valve in the forward direction

3.22

**fracture**

complete separation of any part of the *heart valve repair device* [\(3.25\)](#) ~~(3.25)~~ that was previously intact

3.23

**harm**

injury or damage to the health of people, or damage to ~~the~~ property or the environment

[SOURCE: ISO/IEC Guide 51:2014, 3.1]

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### 3.24

#### hazard

potential source of *harm* (3.23)

[SOURCE: ISO/IEC Guide 51:2014, 3.2]

### 3.25

#### heart valve repair device

*implant* (3.28(3.28)) intended to improve the function of native human heart valves by acting either on the valve apparatus or on the adjacent anatomy (e.g. ventricle, coronary sinus)

Note 1 to entry: See examples of *heart valve repair devices* in Annex B Annex B.

### 3.26

#### heart valve repair system

set of elements provided to repair the native heart valve, consisting of the *heart valve repair device* (3.25(3.25)), *delivery system* (3.12(3.12)), *accessories* (3.1(3.1)) as applicable, packaging, labelling, and instructions

### 3.27

#### imaging modality

method used to visualize and assess native anatomy and/or device position, geometry and/or function

### 3.28

#### implant

device placed surgically or non-surgically into the human body and intended to remain in place after the procedure

### 3.29

#### implant site

intended location of heart valve repair device (3.25) implantation or deployment

### 3.30

#### indication for use

clinical condition of the patient population that the *heart valve repair device* (3.25) is intended to treat or improve

### 3.31

#### intended use

use of a product or process ~~in accordance with~~ according to the specifications, instructions, and information provided by the manufacturer

### 3.32

#### linearized rate

total number of events divided by the total time under evaluation

Note 1 to entry: Generally, the *linearized* rate is expressed in terms of percent per patient year.

### 3.33

#### loading

process to affix, attach or insert a repair device component onto or into a *delivery system* (3.12(3.12))

### 3.34

#### mean arterial pressure

time-averaged arithmetic mean value of the arterial pressure during one *cycle* (3.9)