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Biological evaluation of medical devices - Part 2: Animal welfare requirements (ISO/DIS 10993-2:2024)

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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 documents 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).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

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.

This document was prepared by Technical Committee ISO/TC 194, *Biological and clinical evaluation of medical devices*, in collaboration with the European Committee for Standardization (CEN) Technical Committee CEN/TC 206, *Biological and clinical evaluation of medical devices*, in accordance with the Agreement on technical cooperation between ISO and CEN (Vienna Agreement).

This fourth edition cancels and replaces the third edition (ISO 10993-2:2022).

The main change is the update of the bibliography entries.

A list of all parts in the ISO 10993 series can be found on the ISO website.

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 <u>www.iso.org/members.html</u>.

Introduction

The goal of the ISO 10993 series is the protection of humans in the context of the use of medical devices.

This document supports the goal of the ISO 10993 series by promoting good science through paying proper regard to maximizing the use of scientifically sound non-animal tests and by ensuring that those animal tests performed to evaluate the biological properties of materials used in medical devices are conducted humanely according to recognized ethical and scientific principles.

The application of such humane experimental techniques, including high standards of animal care and accommodation, both help to ensure the scientific validity of safety testing and enhance the welfare of the animals used.

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Biological evaluation of medical devices -

Part 2: Animal welfare requirements

1 Scope

This document specifies the minimum requirements to be satisfied to ensure and demonstrate that proper provision has been made for the welfare of animals used in animal tests to assess the biocompatibility of materials used in medical devices. It is aimed at those who commission, design and perform tests or evaluate data from animal tests undertaken to assess the biocompatibility of materials intended for use in medical devices, or that of the medical devices themselves.

This document makes recommendations and offers guidance intended to facilitate future further reductions in the overall number of animals used, refinement of test methods to reduce or eliminate pain or distress in animals, and the replacement of animal tests by other scientifically valid means not requiring animal tests.

This document applies to tests performed on living vertebrate animals, other than man, to establish the biocompatibility of materials or medical devices.

This document does not apply to tests performed on invertebrate animals and other lower forms; nor (other than with respect to provisions relating to species, source, health status, and care and accommodation) does it apply to testing performed on isolated tissues and organs taken from vertebrate animals that have been euthanized.

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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

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 10993-1 and the following apply.

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

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

3.1

alternative method

test method that replaces an *animal test* (3.3), reduces the numbers of *animals* (3.2) used or refines the procedures applied

3.2

animal

live non-human vertebrate, excluding immature forms during the first half of gestation or incubation

3.3 animal test

use of an *animal* (<u>3.2</u>) for scientific purposes

Note 1 to entry: The definition of an animal test excludes acts of recognized veterinary practice applied for the benefit of an animal or the group of animals of which it is part; recognized husbandry practices to manage or conserve the animal or the group of which it is part; marking by methods which cause no more than momentary pain or distress; and *euthanasia* (3.5).

Note 2 to entry: The prevention of pain, suffering, distress or lasting harm by the effective use of anaesthesia or analgesia or other methods of rendering the animal insentient to pain (e.g. decerebration) does not place animal tests outside the scope of this definition. The administration of anaesthetics, analgesics or other methods of rendering the animal insentient to pain are considered to constitute an integral part of the animal test.

3.4

competent authority

body designated or recognized by a national government to take responsibility for overseeing, supervising or regulating *animal tests* (3.3), or the breeding and supply of *purpose-bred animals* (3.10) for use on such tests, within the scope of this document

3.5

euthanasia

humane killing of an *animal* (3.2) by a method causing minimal physical and mental suffering

3.6

humane endpoint

pre-determined, specific criteria and measures to be implemented to minimize or terminate pain, suffering or distress caused by *animal tests* (3.3) as soon as **a second part of a second par**

the scientific objectives have been met, or

- when it is realized they cannot be met, or
- when the *animal* (<u>3.2</u>) welfare problems being encountered are greater than can be justified by the importance, potential benefits, objectives and nature of the study

3.7

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ps laboratory animal veterinarian ds/sist/bfecbe51-ab9d-4708-b194-036629de5185/osist-pren-iso-10993-2-2025 qualified laboratory animal veterinarian

person responsible for the health and well-being of all laboratory animals (3.2) used at the institution

Note 1 to entry: Often called "attending veterinarian" who is certified or has training or experience in laboratory animal science and medicine or is otherwise qualified in the care of the species being used.

Note 2 to entry: It is recommended that a laboratory animal veterinarian appropriately qualified by the relevant competent authority should be used as an attending veterinarian.

Note 3 to entry: There is an International Association of Colleges of Laboratory Animal Medicine (IACLAM) that is an association of associations, specifically the member Colleges of laboratory animal medicine. Each college has members that, in addition to their demonstrated proficiency in laboratory animal medicine, also possess subspecialization in a variety of areas that have direct bearing on the care, use and welfare of laboratory animals.

3.8

procedural training

prior training and acclimatizing of *animals* (3.2) to the interventions to be performed during an animal test, with a view to minimizing stress to the animal when animal tests are conducted

3.9

protocol

documentation prepared in advance of *animal tests* ($\underline{3.3}$) being undertaken setting out the justification, rationale and test method [including scientific and *humane endpoints* ($\underline{3.6}$)] for the animal tests