Biological evaluation of medical devices —

Part 2: Animal welfare requirements

The European Standard EN ISO 10993-2:1998 has the status of a British Standard
National foreword


The UK participation in its preparation was entrusted to Technical Committee CH/26, Biological testing of medical and dental materials and devices, which has the responsibility to:

— aid enquirers to understand the text;
— present to the responsible international/European committee any enquiries on the interpretation, or proposals for change, and keep the UK interests informed;
— monitor related international and European developments and promulgate them in the UK.

A list of organizations represented on this committee can be obtained on request to its secretary.

For in vivo tests, attention is drawn to the provisions of the Animals (Scientific Procedures) Act 1986, which regulates the use of animals for experimental and other scientific purposes in the UK. The Act requires that:

— Studies be justified and subjected by the Home Office to a cost benefit analysis.
— Animals are only used when it would not be feasible to use alternative methods, and when the standards of care and accommodation equal or surpass the minimum set out in Codes of Practice issued under the Act.
— Protocols of minimum severity are used.

Cross-references

Attention is drawn to the fact that CEN and CENELEC standards normally include an annex which lists normative references to international publications with their corresponding European publications. The British Standards which implement international or European publications referred to in this document may be found in the BSI Standards Catalogue under the section entitled “International Standards Correspondence Index”, or by using the “Find” facility of the BSI Standards Electronic Catalogue.

A British Standard does not purport to include all the necessary provisions of a contract. Users of British Standards are responsible for their correct application.

Compliance with a British Standard does not of itself confer immunity from legal obligations.

Summary of pages

This document comprises a front cover, an inside front cover, pages i and ii, the EN ISO title page, page 2, the ISO title page, pages ii to iv, pages 1 to 4 and a back cover.

This standard has been updated (see copyright date) and may have had amendments incorporated. This will be indicated in the amendment table on the inside front cover.

Amendments issued since publication

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This British Standard, having been prepared under the direction of the Health and Environment Sector Board, was published under the authority of the Standards Board and comes into effect on 15 April 1998.

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Biological evaluation of medical devices — Part 2: Animal welfare requirements

(ISO 10993-2:1992)
Foreword

The text of the International Standard from Technical Committee ISO/TC 194 “Biological evaluation of medical devices” of the International Organization for Standardization (ISO) has been taken over as a European Standard by Technical Committee CEN/TC 206 “Biocompatibility of medical and dental materials and devices”, the secretariat of which is held by NNI.

ISO 10993 consists of the following parts, under the general title “Biological evaluation of medical devices”:

— Part 1: Guidance on selection of tests;
— Part 2: Animal welfare requirements;
— Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity;
— Part 4: Selection of tests for interactions with blood;
— Part 5: Tests for cytotoxicity: in vitro methods;
— Part 6: Tests for local effects after implantation;
— Part 7: Ethylene oxide sterilization residuals;
— Part 9: Degradation of materials related to biological testing;
— Part 10: Tests for irritation and sensitization;
— Part 11: Tests for systemic toxicity;
— Part 12: Sample preparation and reference materials;
— Part 13: Identification and quantification of degradation products from polymers;
— Part 14: Identification and quantification of degradation products from ceramics;
— Part 15: Identification and quantification of degradation products from coated and uncoated metals and alloys;
— Part 16: Toxicokinetic study design for degradation products and leachables;
— Part 17: Glutaraldehyde and formaldehyde residues in industrially sterilized medical devices.

Future parts will deal with other relevant aspects of biological testing.

Annex A of this part of ISO 10993 is for information only.

This European Standard shall be given the status of a national standard, either by publication of an identical text or by endorsement, at the latest by August 1998, and conflicting national standards shall be withdrawn at the latest by August 1998.

According to the CEN/CENELEC Internal Regulations, the national standards organizations of the following countries are bound to implement this European Standard: Austria, Belgium, Czech Republic, Denmark, Finland, France, Germany, Greece, Iceland, Ireland, Italy, Luxembourg, Netherlands, Norway, Portugal, Spain, Sweden, Switzerland and the United Kingdom.

Endorsement notice

The text of the International Standard ISO 10993-2:1992 was approved by CEN as a European Standard without any modification.

NOTE Normative references to International Standards are listed in Annex ZA (normative).
Biological evaluation of medical devices —
Part 2:
Animal welfare requirements

Évaluation biologique des dispositifs médicaux —
Partie 2: Exigences concernant la protection des animaux
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Descriptors: Medical equipment, surgical equipment, surgical implants, dental equipment, tests, biological tests, laboratory animals.
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.

Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75% of the member bodies casting a vote.

International Standard ISO 10993-2 was prepared by Technical Committee ISO/TC 194, Biological evaluation of medical devices.

ISO 10993 consists of the following parts, under the general title Biological evaluation of medical devices:

— Part 1: Guidance on selection of tests;
— Part 2: Animal welfare requirements;
— Part 3: Tests for genotoxicity, carcinogenicity and reproductive toxicity;
— Part 4: Selection of tests for interactions with blood;
— Part 5: Tests for cytotoxicity: in vitro methods;
— Part 6: Tests for local effects after implantation;
— Part 7: Ethylene oxide sterilization residuals;
— Part 8: Clinical investigation;
— Part 9: Degradation of materials related to biological testing;
— Part 10: Tests for irritation and sensitization;
— Part 11: Tests for systemic toxicity;

Future parts will deal with other relevant aspects of biological testing.

Annex A of this part of ISO 10993 is for information only.
Introduction

The protection of humans is the primary goal of the ISO 10993 series of standards. A second equally important goal is to ensure animal welfare and to minimize the number and exposure of the laboratory animals.

This part of ISO 10993 was developed to ensure the welfare of animals used in biological evaluation testing. Therefore, minimum requirements for the care and use of animals are stated.

A list of international documents concerning the care and handling of animals in biomedical research is given in Annex A for information.

1 Scope

This part of ISO 10993 specifies minimum requirements for the use of animals in biological testing.

This part of ISO 10993 is also intended

a) to establish guidelines which allow the scientist to respect life in general;

b) to reduce the number of animal experiments and the number of animals used in experiments, among other ways by optimization of those performed;

c) to minimize suffering and maintain the quality of life of the animals used in the experiments.

This part of ISO 10993 applies to the experimentation performed on vertebrates. It does not apply to experimentation performed on less differentiated animals; nor does it apply to that part of the experimental work performed on isolated tissues and organs.

This part of ISO 10993 also makes recommendations concerned with the aim of reducing the number of animals used for biocompatibility testing and when possible abolishing animal experiments in this area.

2 Normative reference

The following standard contains provisions which, through reference in this text, constitute provisions of this part of ISO 10993. At the time of publication, the edition indicated was valid. All standards are subject to revision, and parties to agreements based on this part of ISO 10993 are encouraged to investigate the possibility of applying the most recent edition of the standard indicated below. Members of IEC and ISO maintain registers of currently valid International Standards.


3 Definitions

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

3.1 animal

any live non-human vertebrate, excluding foetal or embryonic forms, unless otherwise qualified

3.2 experimental animal

animal used or to be used in experiments

3.3 bred animal

animal specially bred for use in experiments in facilities accredited by, or registered with, the competent authority

3.4 animal experiment

any use of an animal for scientific purposes which may cause it pain, anxiety, suffering, distress or lasting harm, excluding the least painful methods accepted in modern veterinary or laboratory practice (i.e. “humane” methods) of killing or marking an animal

an experiment starts when an animal is first prepared for use and ends when no further observations are to be made for that experiment

NOTE 1 The prevention, elimination and minimization of pain, suffering, distress or lasting harm by the successful use of anaesthesia or analgesia or other methods does not place the use of an animal outside the scope of this definition.

3.5 competent authority

that authority designated by each state as being responsible for supervising the experiments within the scope of this part of ISO 10993

3.6 properly anaesthezized

deprived of sensation by methods of anaesthesia (whether local or general) as effective as those used in good veterinary practice

3.7 humane method of killing

killing of an animal with a minimum of physical and mental suffering

NOTE 2 Appropriate means will vary according to the animal species.

3.8 unnecessary repetition

duplication of the same experiment without scientific need
NOTE 3 If experimental results are properly confirmed, further repetitions are considered unnecessary. This statement does not apply to the necessary controls within an experiment.

4 Requirements
NOTE 4 See Annex A for bibliographical references.

4.1 Sequence of in vitro and in vivo tests
Animal experiments shall not be performed before appropriate in vitro tests, if available, have been carried out.

If the in vitro tests clearly show that the material, device or extract is unsuitable the animal experiment shall not be performed.

4.2 Prevention of unnecessary repetition
Scientists proposing to conduct biological evaluation tests shall make diligent efforts to ascertain that any proposed animal experiments have not been done previously. Scientists conducting biological evaluation tests are encouraged to publish the results of their experiments including negative ones in internationally referenced journals, using keywords that allow identification of relevant animal experiments.

Licensing authorities are to be encouraged to establish specific lines of communication directed toward preventing unnecessary repetition. (See 5.2.)

4.3 Availability of results
It is strongly recommended that the results of appropriately performed and evaluated tests be accepted by all countries.

4.4 Qualification of persons involved
All persons involved in performing animal experiments shall be

a) appropriately qualified;\(^1\)

b) suitably trained in the humane care of the animal species being used;

c) trained in all appropriate legislation;

d) trained in the scientific aspects of the research being conducted.

4.5 Care and handling
Care and handling of the animals shall conform to accepted animal husbandry guidelines. Care and handling of the animals shall prevent distress and pain as far as possible. See Annex A.

4.6 Surgical procedure
All surgical procedures on experimental animals, especially those from which the animals are allowed to recover, shall be performed on properly anaesthetized animals using appropriate aseptic procedures and careful handling of tissues involved.

4.7 Pre-, per- and post-operative care
All surgical procedures on experimental animals from which the animals are allowed to recover shall include appropriate provisions for pre-, per- and post-operative care of the animals in accordance with established veterinary medical and nursing practices.

If pre-, per- and post-operative pain is discerned, it shall be recorded and, unless precluded for scientific reasons, it shall be alleviated through the use of appropriate methods of analgesia or the experiment shall be terminated.

4.8 Planning of experiment
The design of the experiment should be appropriate to meet the desired objectives.

The design of an animal experiment shall be specified in an Experimental Plan. In addition, the investigator shall consider the use of non-invasive or alternative methods of investigation to reduce the number of animals used in the experiment (see 4.9).

The Experimental Plan shall contain the following, as appropriate:

a) details of the statistical methods to be applied before and, if necessary, throughout the entire experiment starting with the design of the Experimental Plan and ending with the Final Report;

b) essential information about the composition of the device or material and about the use of the device or material under investigation;

c) the specific goals and the scientific questions to be investigated in the study;

d) the procedures used to conduct the experiment (which should be appropriate to the device or material under investigation) including:

1) the species and approximate number of animals to be used,

2) the rationale for involving animals, and for the appropriateness of the species and numbers used,

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\(^1\) The personality and attitude play an important role in this respect.
3) the origin of the animal in order to minimize the use of animals not bred for experimentation,
4) a description of the proposed use of the animals,
5) a description of any method of euthanasia to be used.

All control procedures and comparators, whether real, standardized or simulated, shall be specified.

4.9 Reduction of animal experiments
The final intention of ISO 10993 is to forego the need for animal experiments. Toward that goal, the planning of the experiment shall consider the use of the least invasive test methods in an animal and/or the reduction of animal experiments by using less invasive methods in the same animal.

4.10 Evaluation
The evaluation of test results shall be thorough and statistical evaluation shall be performed when required.

4.11 Multiple experiments in same animal
In general an animal shall not be used for more than one experiment in a series. The need to avoid undue suffering in the animals used should take precedence over the need to reduce the number of animals used.

4.12 Methods of euthanasia
Methods of euthanasia employed at the termination of animal experiments shall produce rapid unconsciousness and subsequent death without evidence of pain or distress.

5 Recommendations
Recommendations concerned with the future scientific study to reduce the number of animals used in biological testing, to refine the experimental methods to reduce or eliminate pain in animals, and to replace animal experiments by other means, are given in 5.1 to 5.6.

5.1 Alternative methods
Priority should be given by competent authorities, funding agencies and scientists to the validation and/or development of alternative methods. One way in which this could be achieved is to encourage editors of scientific journals to publish papers which describe alternative methods and negative results.

5.2 Database for prevention of unnecessary repetition
International databases should be established to minimize unnecessary repetition.

5.3 Animal care and handling — International documents
It is strongly recommended that internationally accepted detailed documents be produced and updated regarding the care and handling of experimental animals.

5.4 Reduction in animal usage
It is strongly recommended that authorities require only the minimum possible number of animal experiments to be performed in order to yield meaningful data and not maximum precision.

5.5 Pilot experiments
Pilot experiments should be performed to allow planning of a minimum number of experiments to provide the required result. If in a standard test the minimum number of animals required is given, that number takes precedence.

5.6 Guidelines for animal husbandry
It is requested that documents on updated animal husbandry guidelines be forwarded to ISO/TC 194.

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2) While appreciating the question of confidentiality, it is recommended that this should not preclude the creation of the database.
Annex A (informative)

Bibliography

[10] Law concerning the protection and control of animals (Japan).

NOTE 5 Other documents, in accordance with 5.6, will be added when available.

Annex ZA (normative)

Normative references to international publications with their relevant European publications

This European Standard incorporates by dated or undated reference, provisions from other publications. These normative references are cited at the appropriate places in the text and the publications are listed hereafter. For dated references, subsequent amendments to or revisions of any of these publications apply to this European Standard only when incorporated in it by amendment or revision. For undated references the latest edition of the publication referred to applies.

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