
**Ophthalmic implants — Intraocular
lenses —**

**Part 3:
Mechanical properties and test methods**

*Implants ophtalmiques — Lentilles intraoculaires —
Partie 3: Propriétés mécaniques et méthodes d'essai*
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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.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. 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.

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.

ISO 11979-3 was prepared by Technical Committee ISO/TC 172, *Optics and photonics*, Subcommittee SC 7, *Ophthalmic optics and instruments*.

This second edition cancels and replaces the first edition (ISO 11979-3:1999), which has been technically revised.

ISO 11979 consists of the following parts, under the general title *Ophthalmic implants — Intraocular lenses*:

- *Part 1: Vocabulary* <https://standards.iteh.ai/catalog/standards/sist/5a60a519-88ce-4088-97ad-22ab4714792c/iso-11979-3-2006>
- *Part 2: Optical properties and test methods*
- *Part 3: Mechanical properties and test methods*
- *Part 4: Labelling and information*
- *Part 5: Biocompatibility*
- *Part 6: Shelf-life and transport stability*
- *Part 7: Clinical investigations*
- *Part 8: Fundamental requirements*
- *Part 9: Multifocal intraocular lenses*
- *Part 10: Phakic intraocular lenses*

Introduction

This part of ISO 11979 contains methods for which requirements are given and methods for which no requirements are formulated. The former are considered essential for the safety or performance of the intraocular lens, while the latter provide essential information to the ophthalmic surgeon or are used for other purposes.

A special purpose is the use of mechanical data to assess the need for clinical investigation of modifications of existing models as described in ISO 11979-7 [7]. Because of the complexity of this analysis, detailed descriptions and examples have been given in ISO/TR 22979 [8]. Due to the wide variety of intraocular lens designs already on the market, it has not been possible to devise test methods that are applicable to every design under all circumstances. It can be anticipated that new materials currently under development will result in drastically new designs that will require modified or other test methods. As with all standards, it is then up to the parties using the standard to modify or develop corresponding methods, and give rationale and validation for them in a spirit that is consistent with this part of ISO 11979.

In cases where different tolerances have been given depending on material or design, they reflect an existing situation with well-established products.

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Ophthalmic implants — Intraocular lenses —

Part 3: Mechanical properties and test methods

1 Scope

This part of ISO 11979 specifies requirements and test methods for certain mechanical properties of intraocular lenses (IOLs).

It is applicable to all types of IOLs intended for implantation in the anterior segment of the human eye, excluding corneal implants, provided that the test method is appropriate to the particular IOL design.

2 Normative references

The following referenced documents are indispensable for the application 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 11979-1, *Ophthalmic implants — Intraocular lenses — Part 1: Vocabulary*
<https://standards.iteh.ai/catalog/standards/sist/5a60a519-88ce-4088-97ad-27ab4714792e/iso-11979-1-2006>

ISO 11979-2, *Ophthalmic implants — Intraocular lenses — Part 2: Optical properties and test methods*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 11979-1 apply.

4 Requirements

4.1 General

For all IOLs, the mechanical properties shall be determined at *in situ* conditions. The precise composition of the solution used shall be reported in all cases. Alternative test conditions, e.g. room temperature conditions, may be used if a justification to deviate from *in situ* is given. The alternative test conditions shall be specified in the test reports.

For each of the methods described below, tests shall be performed on a minimum of three IOL lots of medium dioptric power. If dioptric power affects the property tested, the lots shall comprise one each of low, medium and high dioptric powers. The minimum sample size for each test shall be 10 IOLs per lot. The lots shall be representative of IOLs being marketed. In all cases, the sampling criteria applied shall be reported. Means and standard deviations shall be reported for the pooled samples.

If, for certain designs and certain applications, a specific test method described in this part of ISO 11979 is not applicable, the IOL manufacturer can devise a corresponding test method and provide a validation and rationale for it.

4.2 Tolerances and dimensions

For all types of IOLs except multipiece posterior chamber IOLs, the tolerance on the overall diameter shall be $\pm 0,20$ mm. For multipiece posterior chamber IOLs, the tolerance on the overall diameter shall be $\pm 0,30$ mm.

The tolerance on the vault height shall be as follows:

- a) for anterior chamber IOLs, $\pm 0,15$ mm;
- b) for multipiece posterior chamber IOLs, $\pm 0,35$ mm;
- c) for other IOLs, $\pm 0,25$ mm.

The tolerance on the sagitta shall be as follows:

- for anterior chamber IOLs, $\pm 0,25$ mm;
- for multipiece posterior chamber IOLs, $\pm 0,45$ mm;
- for other IOLs, $\pm 0,35$ mm.

The tolerance on the clear optic shall be $\pm 0,15$ mm. The diameter of the clear optic shall be greater than 4,25 mm in any meridian. The tolerance on the dimensions of the body shall be $\pm 0,10$ mm. For ellipsoid IOLs, the dimensions of the body shall be reported as (short axis) \times (long axis).

The tolerance on the diameter of the positioning hole shall be nominal $\left(\begin{smallmatrix} +0,05 \\ 0 \end{smallmatrix} \right)$ mm.

Dimensions for which tolerances are given above shall be specified in the manufacturer's design documentation. Some dimensions may vary with dioptric power, hence different specifications may apply to individual powers of an intraocular lens design. [ISO 11979-3:2006](https://standards.iteh.ai/catalog/standards/sist/5a60a519-88ce-4088-97ad-22ab4714792c/iso-11979-3-2006)

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4.3 Clearance analysis

An empirical analysis of anatomic placement shall be performed for anterior chamber lenses to evaluate the most proximate points with relation to the anatomical structures of the eye. The clearance of the anterior surface of the IOL optic in relation to the endothelial layer of the cornea shall be determined for the lens at its minimum recommended diameter in its compressed state. The separation of the posterior surface of the IOL optic to the iris as well as if phakic to the crystalline lens shall be determined. These results shall be considered in the risk analysis. The theoretical eye model in Annex I can be used in the evaluation.

The manufacturer should strive for a clearance of at least 1 mm under worst-case conditions, i.e. conditions which would result in the minimum amount of clearance.

4.4 Compression force

Using the method described in Annex A, the compression force shall be measured and reported as follows:

- a) for IOLs intended for capsular bag placement, with the haptics compressed to a diameter of 10 mm;
- b) for IOLs intended for sulcus placement, with the haptics compressed to a diameter of 11 mm;
- c) for IOLs intended for both capsular bag and sulcus placement, with the haptics compressed to both a diameter of 10 mm and a diameter of 11 mm;
- d) for anterior chamber IOLs, with the haptics compressed to the minimum and maximum intended compressed diameters recommended by the manufacturer in the product literature.

4.5 Axial displacement in compression

Using the method described in Annex B, the axial displacement in compression shall be measured and reported at the same diameters that were used for the measurement of compression force (see 4.4).

In addition, for anterior chamber IOLs, the vault height and the sagitta in the compressed state shall be given in the product literature as a function of dioptric power at the minimum and maximum intended compressed diameters, as specified in 4.4.

4.6 Optic decentration

Using the method described in Annex C, the optic decentration shall be measured and reported at the same diameters that were used for the measurement of compression force (see 4.4).

The sum of the arithmetic mean and two standard deviations of the optic decentration shall not exceed 10 % of the clear optic.

4.7 Optic tilt

Using the method described in Annex D, the optic tilt shall be measured and reported at the same diameters that were used for the measurement of compression force (see 4.4).

The sum of the arithmetic mean and two standard deviations of the optic tilt shall not exceed 5°.

4.8 Angle of contact

Using the method described in Annex E, the angle of contact shall be measured and reported at the same diameters that were used for the measurement of compression force (see 4.4).

NOTE The angle of contact is a measured approximation of the total haptic contact with the supporting ocular tissue.

4.9 Compression force decay

Using the method described in Annex F, the compression force decay shall be tested and reported at the same diameters that were used for the measurement of compression force (see 4.4).

The loops of IOLs are designed to exert some pressure on eye structures as a means of keeping the IOL in position and should continue to do so for some time after implantation.

Results shall be reported as residual compression force after $24 \text{ h} \pm 2 \text{ h}$ in compression at each required compressed diameter.

4.10 Dynamic fatigue durability

All loops shall be capable of withstanding, without breaking, 250 000 cycles of near-sinusoidal deformation of $\pm 0,25 \text{ mm}$ around the compressed distance.

Using the method described in Annex G, fatigue testing shall be performed as follows:

- a) for IOLs intended for capsular bag placement, at a compressed distance of 5,0 mm between the testing plate and the centre of the optic;
- b) for IOLs intended for sulcus placement, at a compressed distance of 5,5 mm between the testing plate and the centre of the optic;
- c) for IOLs intended for both capsular bag and sulcus placement, at a compressed distance of 5,0 mm between the testing plate and the centre of the optic;

- d) for anterior chamber IOLs, at a distance between the testing plate and the centre of the optic, corresponding to half the maximum intended compressed diameter as recommended by the manufacturer in the product literature.

This test shall be carried out only for IOL designs in which the loop will be in a compressed state when implanted. The frequency shall be between 1 Hz and 10 Hz.

Higher frequencies can be used if it is verified that the loop follows the testing plate without lag at all times.

No loop tested shall break.

For IOLs designed to move axially under compression, additional testing shall be considered.

4.11 Surgical manipulation

The IOL manufacturer shall provide evidence that the loops of an IOL design are capable of withstanding surgical manipulations without failure. An appropriate test method and specification shall be established by the manufacturer to ensure that the device does not fail at typical deformations. A test method, useful for some designs with loops, is given in Annex H.

4.12 Surface and bulk homogeneity

The IOL shall be essentially free from defects, i.e. deviations from surface and bulk homogeneity that are not intended features of the design, including all kinds of surface defects such as scratches, digs, protrusions, cracks, roughness, etc., as well as bulk defects such as inclusions, bubbles, striae, discoloration, etc. The lens should be inspected at 10× under optimal lighting conditions; any questionable or critical areas should be viewed at higher magnification.

5 Recovery of properties following simulated surgical manipulation

The testing in this clause applies only to IOLs of which the optic is intended to be folded or compressed during implantation. Perform testing on 10 lenses of each of the dioptric powers with the smallest and largest cross-sectional dimensions. In practice this will typically be 10 lenses with the lowest and 10 lenses with the highest dioptric power. Follow the instructions supplied by the manufacturer, using recommended lubricants and instrumentation. To determine the acceptable time during which the lens is allowed to be kept deformed prior to implantation, maintain the deformed state for a period of time. This time shall not be shorter than 3 min. Times in excess of 20 min need not be investigated. The time used shall be reported.

After release from the deformed state, allow the lens to relax at *in situ* conditions up to 24 h ± 2 h. The time used shall be reported. Subsequently:

- a) measure dioptric power and image quality (see ISO 11979-2);
- b) measure overall diameter and sagitta (see 4.2);
- c) inspect for surface and bulk homogeneity (see 4.12).

The results shall be reported and are acceptable if they remain within manufacturing specifications of the product.

Annex A (normative)

Measurement of compression force

A.1 Principle

The force exerted by the loops is measured when the IOL is confined to a prescribed diameter with the movement of the body being unrestricted.

A.2 Apparatus

A diagram of the apparatus is shown in Figures A.1 and A.2 and comprises the following.

A.2.1 Two anvils, with faces having a radius of $5,00 \text{ mm} \pm 0,02 \text{ mm}$ or $5,50 \text{ mm} \pm 0,02 \text{ mm}$, as appropriate, constructed from a low-friction material to minimize loop rotational constraint, and aligned relatively with each other.

A.2.2 Device, capable of measuring force accurate to at least $\pm 0,1 \text{ mN}$.

A.3 Procedure

A.3.1 Carry out the testing with the IOL in the horizontal plane.

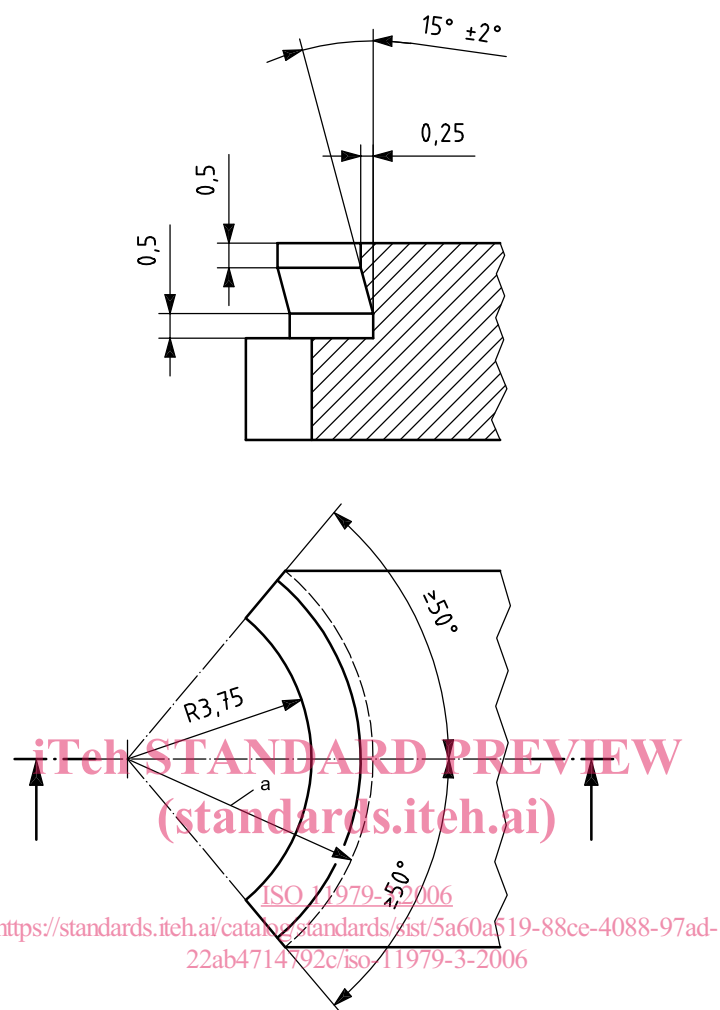
NOTE Testing in the vertical plane leads to asymmetrical distribution of force between the loops due to the mass of the IOL.

A.3.2 Set the anvils to a distance approximately equal to the overall dimension of the IOL and place the IOL between the anvils.

A.3.3 Locate the IOL in the uncompressed state so that the line of compression bisects the angle of contact in the compressed state or, in the case of IOLs where there are multiple contacts, so that the line of compression bisects the angle of contact of the extremes in the compressed state (see Figure A.3).

A.3.4 Close the anvils to the prescribed diameter.

A.3.5 Read the compression force after allowing between 10 s and 30 s for the IOL to stabilize.



^a $R = 5,00 \text{ mm} \pm 0,02 \text{ mm}$ or $5,50 \text{ mm} \pm 0,02 \text{ mm}$.

Figure A.1 — Anvil