
**Ophthalmic optics — Contact lenses and
contact lens care products — Guidance
for clinical investigations**

*Optique ophtalmique — Lentilles de contact et produits d'entretien pour
lentilles de contact — Directives pour les investigations cliniques*

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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 11980 was prepared by Technical Committee ISO/TC 172, *Optics and photonics*, Subcommittee SC 7, *Ophthalmic optics and instruments*.

This third edition cancels and replaces the second edition (ISO 11980:2009), which has undergone minor revision in order to update the normative reference to ISO 14155 and to revise 4.2.1.1 b) 6) and the fifth row of Table A.1 (overnight wear).

This corrected version of ISO 11980:2012 incorporates the following correction:

- in Table A.12, the final equation corresponding to the total number of eyes has been inserted.

Introduction

Currently, contact lenses and contact lens care products are regulated in different ways in different countries. This International Standard has been developed to encourage global harmonization. Widespread adoption of this International Standard should represent yet another step toward mutual recognition. This International Standard can also be used as a basis to fulfil design elements of ISO 9001^[1].

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Ophthalmic optics — Contact lenses and contact lens care products — Guidance for clinical investigations

1 Scope

This International Standard gives guidelines for the clinical investigation (CI) of the safety and performance of contact lenses and contact lens care products.

NOTE This International Standard attempts to harmonize the recognized regulatory requirements for the conduct of a CI to meet the marketing and labelling requirements for contact lenses and contact lens care products around the world. However, national requirements vary greatly. Wherever national practice or regulations dictate some legal requirement, this requirement takes precedence over this International Standard.

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 14155, *Clinical investigation of medical devices for human subjects — Good clinical practice*

ISO 14534, *Ophthalmic optics — Contact lenses and contact lens care products — Fundamental requirements*

ISO 18369-1, *Ophthalmic optics — Contact lenses — Part 1: Vocabulary, classification system and recommendations for labelling specifications*

3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 14155, ISO 14534 and ISO 18369-1 apply.

4 Clinical investigational requirements

4.1 General

The general requirements for a CI and for a clinical investigation plan (CIP) given in ISO 14155 shall apply, with additional requirements given below.

4.2 Additional requirements

4.2.1 Study design

4.2.1.1 General

- a) The inclusion criteria for subject selection shall relate to the study objectives and should include:
- 1) subjects with normal eyes who are not using any ocular medications, aged 18 years or over [except when contact lens investigations have a special indication for use in “children” (for the purposes of this International Standard, persons less than 18 years of age) such as orthokeratology and paediatric aphakic lenses];
 - 2) lens powers within the range available for the test lenses;
 - 3) the manifest cylinder less than or equal to 0,75 D (for a study with only spherical power correcting lenses);

- 4) best spectacle corrected visual acuity greater than or equal to 20/25 (less than or equal to LogMAR 0,1).
- b) The exclusion criteria for subject selection shall relate to the study objectives and should include, but not be limited to:
 - 1) anterior segment infection, inflammation or abnormality;
 - 2) any active anterior segment ocular disease that would contraindicate contact lens wear;
 - 3) the use of systemic or ocular medications that would contraindicate contact lens wear;
 - 4) history of herpetic keratitis;
 - 5) history of refractive surgery or irregular cornea (except when the contact lenses under investigation are indicated for irregular cornea, keratoconus or refractive surgery);
 - 6) slit lamp findings that are more serious than grade 1;
 - 7) corneal vascularization greater than 1 mm of penetration;
 - 8) a pathologically dry eye;
 - 9) participation of the subject in a contact lens or contact lens care product clinical trial within the previous 30 days.
- c) The CIP shall provide a description of the monitoring procedure to ensure consistent quality of data collection and recording.
- d) The CIP shall include a statistical analysis plan. Sample size shall be justified, calculated by a validated statistical software package.

4.2.1.2 Contact lenses

4.2.1.2.1 General. A CI of contact lenses, including daily wear and extended wear hydrogel, silicone hydrogel, and rigid gas-permeable contact lenses, shall be designed as one of 4.2.1.2.2 or 4.2.1.2.3.

For CIPs to demonstrate safety and performance, as well as special claims (e.g. comfort), labelling or additional indications, the following is required: a pre-determined statistical analysis plan (including sample size calculations) shall be specified in the clinical protocol. Where feasible, the CIP shall define objective endpoints to help support such claims.

NOTE 1 Inter-subject controls are preferred to intra-subject controls due to the potential dependence between the two eyes and concerns regarding subject compliance.

NOTE 2 Annex A provides guidance for the design of a CI.

4.2.1.2.2 As a prospective, concurrently controlled study. For investigations evaluating hydrogel, silicone hydrogel or rigid gas-permeable contact lenses, a prospective, concurrent control study design shall be followed. Either a bilateral crossover design or a contra-lateral eye (i.e. intra-subject) design or inter-subject controls shall be utilized. If inter-subject controls are utilized, the ratio of test subjects to control subjects may be either 2:1 or 1:1. The control lens shall be a currently marketed contact lens in use for the same modality. Randomization and masking (subject, investigator and evaluator) shall be employed where possible to minimize the potential for bias. Subjects shall be divided evenly between study investigators.

4.2.1.2.3 As an uncontrolled study. Here, results are compared to a historical control. Alternative investigational study designs, such as historical controls, shall be utilized when a sponsor has a clinical database on a marketed contact lens to use as a comparator. If any historical control is used, the control group shall be defined and adequately characterized for comparison to the test group. Compatibility of test and control groups shall be demonstrated by comparison of the selection criteria, demographics, refractive characteristics, contact lens wearing history and CIPs used.

4.2.1.3 Contact lens care products

For investigations evaluating contact lens care products, a prospective concurrent control study design shall be followed. It is recommended that the ratio of test to control subjects be either 2:1 or 1:1. The control care product shall be a currently marketed contact lens care product. Randomization and masking (subject, investigator and evaluator) shall be employed where possible to minimize the potential for bias. Subjects shall be divided evenly between study investigators. Alternative investigational study designs, such as use of historical controls, may be utilized when a manufacturer has a clinical database on a marketed care product to use for comparison. If any historical control is used, the control group should be defined and adequately characterized for comparison to the test group. Compatibility of test and control groups should be demonstrated by comparison of the selection criteria and CIPs used.

For CIPs to demonstrate safety and performance, as well as special claims (e.g. comfort), labelling or additional indications, the following is required for the care products: a pre-determined statistical analysis plan (including sample size calculations) shall be specified in the clinical protocol. Where feasible, the protocol should define objective endpoints to help support such claims.

NOTE 1 Inter-subject controls are preferred to intra-subject controls due to the potential dependence between the two eyes and concerns regarding subject compliance.

In a contact lens care product investigation, a daily wear schedule shall be followed for most products in order to maximize the subject's exposure to those products. However, a study of a lens or a periodic cleaner, used at weekly intervals, may provide more valuable clinical data concerning efficacy when extended wear subjects are enrolled than a similar investigation with daily wear subjects.

When a daily wear schedule is used and safety is a primary objective, one post-dispensing visit should be done 1 h to 2 h after lens insertion in order to permit observation of corneal and conjunctival staining caused by an immediate toxicity reaction.

A contact lens care product with a cleaning indication shall have an objective measure of lens cleanliness on at least one lens collected from each subject at the end of the clinical study.

If the manufacturer of a contact lens care product wishes to recommend its use with a specific type of lens in the labelling, the compatibility with the lens type should be confirmed pre-clinically and during the clinical trial.

If the CI has not collected any data on use with a particular type of lens material (such as silicone hydrogel lenses), the product label should clearly state this fact.

NOTE 2 Annex A provides guidance for the design of a CI.

4.2.2 Variables

4.2.2.1 Contact lenses

The following variables should be considered during the CI for contact lenses, in addition to the variables listed in 4.2.2.2:

- a) visual performance;
- b) refractive performance;
- c) keratometric measurements;
- d) lens centration;
- e) lens movement;
- f) lens surface wettability;
- g) lens surface deposits;
- h) subject acceptance of comfort;