# INTERNATIONAL STANDARD

ISO 11979-7

Second edition 2006-05-01

# Ophthalmic implants — Intraocular lenses —

Part 7: Clinical investigations

Implants ophtalmiques Lentilles intraoculaires —
Partie 7: Investigations cliniques

Citck to view the citch to view the citck to view the citch to view the



#### PDF disclaimer

This PDF file may contain embedded typefaces. In accordance with Adobe's licensing policy, this file may be printed or viewed but shall not be edited unless the typefaces which are embedded are licensed to and installed on the computer performing the editing. In downloading this file, parties accept therein the responsibility of not infringing Adobe's licensing policy. The ISO Central Secretariat accepts no liability in this area.

Adobe is a trademark of Adobe Systems Incorporated.

Details of the software products used to create this PDF file can be found in the General Info relative to the file; the PDF-creation parameters were optimized for printing. Every care has been taken to ensure that the file is suitable for use by ISO member bodies. In the unlikely event that a problem relating to it is found, please inform the Central Secretariat at the address given below.

STANDARDSISO COM. Click to view the full POF of 150 Meta T. 2006

#### © ISO 2006

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office Case postale 56 • CH-1211 Geneva 20 Tel. + 41 22 749 01 11 Fax + 41 22 749 09 47 E-mail copyright@iso.org Web www.iso.org

Published in Switzerland

Cont	tents	Page
Forewo	ord	iv
1	Scope	
2	Normative references	1
3	Terms and definitions	1
4	Justification for a clinical investigation	1
5	Ethical considerations	1
6	Terms and definitions  Justification for a clinical investigation  Ethical considerations  General requirements  General	2
6.1 6.2	GeneralAdditional requirements	2
Annex	Additional requirements  A (informative) Elements of a clinical investigation	5
Annex	c Β (informative) Evaluation of post-operative adverse event a	id visual acuity rates13
Bibliog	graphy	18
Ö	graphy	

## **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 11979-7 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 11979-7:2001), which has been technically revised.

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

- Part 1: Vocabulary
- 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

# Ophthalmic implants — Intraocular lenses —

# Part 7:

# **Clinical investigations**

# 1 Scope

This part of ISO 11979 specifies particular requirements for clinical investigation for posterior and anterior chamber monofocal intraocular lenses (IOLs) for the correction of aphakia.

# 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

ISO 14155-1, Clinical investigation of medical devices for human subjects — Part 1: General requirements

ISO 14155-2, Clinical investigation of medical devices for human subjects — Part 2: Clinical investigation plans

## 3 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 11979-1, ISO 14155-1 and ISO 14155-2 apply.

# 4 Justification for a clinical investigation

The requirements given in ISO 14155-1 shall apply.

If a new model is a minor modification of a model for which the safety and performance have been established through clinical investigation in accordance with this part of ISO 11979, no or limited clinical investigation is needed. ISO/TR 22979 provides guidance in determining if a modification is minor.

## 5 Ethical considerations

For clinical investigations of medical devices for human subjects, the requirements in ISO 14155-1 shall apply.

# General requirements

#### General 6.1

The general requirements for a clinical investigation given in ISO 14155-1 and the clinical investigation plan requirements in ISO 14155-2 shall apply, with additional requirements given below.

#### Additional requirements 6.2

## 6.2.1 Design

A clinical investigation of an IOL model shall be designed in one of two ways:

- as an uncontrolled study, in which case the results are compared to the adverse events and visual acuity rates given in Annex B.
- as a controlled study, with the provision that the statistical power to detect differences in the adverse Selso. Com. Circk to view the full PDF of 1 event rates and visual acuity is similar to the uncontrolled study. The controllens shall conform with applicable parts of ISO 11979.

NOTE Annex A provides guidance for the design of a clinical investigation.

#### 6.2.2 Variables

The following variables shall be considered:

- best spectacle corrected visual acuity (BSCVA);
- refraction;
- intraocular pressure;
- corneal status;
- iritis;
- IOL decentration;
- IOL tilt:
- IOL discoloration
- IOL opacity
- cystoid macular oedema;
- hypopyon;
- endophthalmitis;
- pupillary block;
- retinal detachment;
- status of anterior and posterior capsule.

Additional variables can be studied in the clinical investigation to support specific claims.

#### 6.2.3 Other considerations

To minimize the risks associated with the clinical investigation of a new IOL, subject enrolment shall occur in stages. The subject data from each stage shall be evaluated and found acceptable by the sponsor and the coordinating investigator prior to the continuation of the clinical investigation. Guidance on phased enrolment is included in Annex A.

Only the first eye of each subject shall be included in the primary statistical analysis.

Any plans for fellow eye implantation shall be described in the clinical investigation plan. Bilateral implantation shall not be implemented until initial safety and effectiveness data have been collected, evaluated and confirmed by the sponsor and principal investigators.

The review of data from at least 50 eyes with six months of follow-up is recommended Previous clinical experience, i.e. results from well-documented clinical investigations, may be adequate justification to begin bilateral implantation earlier in the study.

The duration of the clinical investigation shall be one year for all posterior chamber IOLs, and 3 years for all anterior chamber IOLs.

The clinical investigation plan shall contain descriptions of the surgical technique, the intraoperative use of ophthalmic viscosurgical devices, and the use of preoperative, intra-operative and post-operative medications. Any deviation shall be recorded on the case report forms.

The clinical investigation plan shall describe how subject visits and ophthalmic adverse events in between reporting periods will be handled in the data analyses.

All subjects in a clinical investigation shall be monitored for the duration of the investigation. The clinical investigation shall be considered completed when all subjects that have been enrolled in the investigation, including subjects whose IOL was removed or replaced, have reached the final reporting period.

Serious ophthalmic adverse events and all adverse device effects shall be reported using a special case report form and forwarded to the sponsor as required. All other ophthalmic adverse events shall be reported using the standard visit case report forms and are collected during monitoring.

# Annex A

(informative)

# Elements of a clinical investigation

#### A.1 General

The following are elements of a clinical investigation plan which can assist in collecting data for the purpose of determining the safety and performance of IOLs.

NOTE This annex reflects the experience with clinical investigations of IOLs in the USA.

# A.2 Number of subjects

The clinical investigation includes a minimum of 300 subjects when the results are compared to the safety and performance endpoints in Annex B. In the case of a study with a concurrent control group, calculate the number of subjects sufficient to detect differences in the safety and performance endpoints in Annex B with similar statistical power to the study mentioned above. Any additional claims, beyond those for safety and performance, require calculation of a sample size for that purpose.

To take into account that some subjects are lost during the course of the clinical investigation (including deceased subjects and subjects who have the IOL explanted), enrol about:

- a) 390 subjects in the one-year investigation;
- b) 500 subjects in the three-year investigation.

Significantly larger numbers of subjects are not to be enrolled in order to minimize exposure to the risks of a new IOL.

To assist in achieving a balance in the number of subjects from each investigator, each surgeon contributes a minimum of 20 subjects, but no more than 25 % of the subjects in the investigation.

If the risk analysis determines that a limited clinical investigation is sufficient (see ISO/TR 22979), then enrol 125 subjects.

# A.3 Phased enrolment

To minimize the potential risks, the clinical investigation consists of two phases as follows.

#### a) Phase 1:

A maximum of 100 subjects are included. After at least 50 of those have reached case report Form 4, their data are evaluated. If the results are acceptable, the next phase can begin.

#### b) **Phase 2**:

The remainder of the subjects are included.

# A.4 Reporting periods

The time frames for the reporting periods are defined below:

- a) Case report Form 0: pre-operative/operative reporting;
- b) Case report Form 1: post-operative reporting 1 d or 2 d post-operatively;
- c) Case report Form 2: post-operative reporting 7 d to 14 d post-operatively;
- d) Case report Form 3: post-operative reporting 30 d to 60 d post-operatively;
- e) Case report Form 4: post-operative reporting 120 d to 180 d post-operatively;
- f) Case report Form 5: post-operative reporting 330 d to 420 d post-operatively;
- g) Case report Form 6: post-operative reporting 630 d to 780 d post-operatively;
- h) Case report Form 7: post-operative reporting 990 d to 1 140 d post-operatively.

The minimum number of completed case report forms for each reporting period is 300.

# A.5 Standardization of the clinical evaluation

Define criteria for evaluation of all studied variables. Define testing conditions for all measurements. Before commencing the investigation instruct and train all investigators to use these, in order to obtain data that can be combined for the purpose of statistical analysis.

# A.6 Data analysis

Consider the following analyses:

- a) VA stratified by age;
- b) best-case VA;
- c) VA stratified by adverse event;
- d) VA stratified by pre-operative ocular pathology;
- e) VA stratified by investigator;
- f) subject-by-subject analysis of reasons why subject failed to achieve 0,5 (6/12; 20/40) VA;
- g) rate of visual acuity decrease of 10 letters or more on an early treatment of diabetic retinopathy study (EDTRS) chart (or equivalent) between a form evaluation and a later form evaluation with the cause of the visual acuity decrease described in each case;
- h) rates of cumulative adverse events stratified by age;
- i) rates of persistent adverse events stratified by age;
- j) adverse event stratified by investigator.

# A.7 Subject accountability

The general requirement for accountability of subjects is given in ISO 14155-1. More specific guidance for subject accountability at each of the post-operative visits in IOL clinical investigations is provided in Table A.1.

Table A.1 — Accountability by post-operative visit

Subject status	Total number					
Subject status	Enrolled a	Form 1	Form 2, etc.	Final form		
	$N_{tot}$					
Available for analysis <sup>b</sup> , n <sub>aa</sub>		$n_{\rm aa}$ , $(n_{\rm aa}/N_{ m tot})$ %	$n_{\rm aa}, \ (n_{\rm aa}/N_{ m tot})$ %	$n_{aa}$ , $(n_{aa}/N_{tot})$ %.		
Missing subjects:				101		
Discontinued <sup>c</sup> , n <sub>d</sub>		$n_{d}$ , $(n_{d}/N_{tot})$ %	$n_{\rm d}$ , $(n_{\rm d}/N_{\rm tot})$ %	$n_{\rm d}$ , $(n_{\rm d}/N_{\rm tot})$ %		
Missing at scheduled visit but seen later $^{\rm d}$ , $n_{\rm sl}$		$n_{\rm sl}$ , $(n_{\rm sl}/N_{\rm tot})$ %	$(n_{\rm sl}/N_{\rm tot})$ %	$n_{\rm sl}, (n_{\rm sl}/N_{\rm tot}) \%$		
Not seen but accounted for $^{\rm e}$ , $n_{\rm ns}$		$n_{\rm ns}$ , $(n_{\rm ns}/N_{\rm tot})$ %	$n_{\rm ns}$ , $(n_{\rm ns}/N_{\rm tot})$ %	$n_{\rm ns}$ , $(n_{\rm ns}/N_{\rm tot})$ %		
Lost to follow-up $^{f},n_{lf}$		$n_{\rm lf}$ $(n_{\rm lf}/N_{ m tot})$ %	$n_{\rm lf}$ , $(n_{\rm lf}/N_{\rm tot})$ %	$n_{\rm lf}$ , $(n_{\rm lf}/N_{\rm tot})$ %		
Active <sup>g</sup> , n <sub>a</sub>	. 0,5	$n_{\rm a}$ , $(n_{\rm a}/N_{\rm tot})$ %	$n_{a}$ , $(n_{a}/N_{tot})$ %	$n_{a}$ , $(n_{a}/N_{tot})$ %		

Explanation of symbols:

n represents the number of subjects associated with the form for that type of information.

 $(n/N_{\text{tot}})$  % represents the percentage of subjects associated with the form of that type of information with respect to the total number of subjects enrolled in the study.

- <sup>a</sup> "Enrolled" or  $N_{\text{tot}}$  represents the total number of subjects enrolled in the investigation.
- $^{\rm b}$  "Available for analysis" or  $n_{\rm aa}$  represents the total number of subjects for whom data is available at the form.
- $^{\rm C}$  "Discontinued" or  $n_{\rm d}$  represents the total number of subjects that have discontinued treatment prior to the form for any reason (e.g. death or device replacement). This category doesn't include subjects that are lost to follow-up.
- $^{
  m d}$  "Missing at final scheduled visit but seen later" or  $n_{
  m Sl}$  represents the total number of subjects that were seen outside the time window associated with the form.
- <sup>e</sup> "Not seen but accounted for" or  $n_{\text{ns}}$  represents the total number of subjects that were missing at the scheduled visit but were accounted for by being contacted (e.g. by phone).
- f c"Lost to follow-up" or  $n_{\rm lf}$  represents the total number of subjects that have missed the form and there is no information available about them.
- <sup>g</sup> "Active" or  $n_a$  represents the total number of subjects that have not reached the time associated with the form. The investigation at the form is considered completed when the number of active subjects is zero.

The following equation is used to determine the percent accountability,  $N_{account}$ , for the investigation.

$$% N_{\text{account}} = \frac{n_{\text{aa}}}{N_{\text{tot}} - n_{\text{d}} - n_{\text{a}}}$$

where  $n_{\rm aa}$ ,  $N_{\rm tot}$ ,  $n_{\rm d}$  and  $n_{\rm a}$  are as defined in Table A.1.

Depending upon the clinical investigation, the total number of subjects is not necessarily the total number of eyes. For the purposes of this guidance, it is assumed that treatment is unilateral and that the total number of subjects is equivalent to the total number of eyes.

To minimize the uncertainty in the data, the lost-to-follow-up subjects in the three-year investigation should be less than 30 % and the lost-to-follow-up in one-year investigation should be less than 10 %.

# A.8 Clinical case report forms

The next pages provide examples of the following case report forms:

- common co c) pre-operative/operative case report form — anterior chamber lenses (Table A.5): Operative case report form — anterior chamber lenses (Table A.5): Operative case report form — anterior chamber lenses (Table A.5): Operative case report form (Table A.6).

  e) adverse event case report form (Table A.6).

  Circle to vice where the chamber lenses (Table A.5): Operative case report form (Table A.6).

Table A.2 — Pre-operative/ operative case report form for posterior chamber lens clinical investigation

Investigator name:			Clinical trial number:	
Patient number: Patient	initials:		Sex: Male:   Race: Caucasian	
			Female: □ Black □ Asian □	
Date of birth:			Other 🗆	
YY MM DD			Mixed $\square$	
Pre-operative report			Irrigating solution used yes no	)
	TY MM			ı
Operative eye	right □	left □	If yes, specify	
	Operative eye	Fellow eye	Periocular medication yes (specify, if appropriate) no	)
Best corrected visual acuity			Anaesthetic	ı
or check one:			Antibiotic	1
Finger count			Corticosteroid	ı
Hand movement			Other (specify)	ı
Light perception			Incision	
No light perception				
IOP (applanation): Op. eye: m	ımHq Fellow eye	: mmHg	Sizemm	
Corneal status (check yes or no for ear		no	Type (e.g., corneal, limbal, scleral tunnel)	
Normal			Type of lens extraction (check one)	
Guttata			Type of total extraordin (stock only)	
Other pathology (specify)	_		Phacoemulsification	
Cataract	_		Other (specify)	
Etiology (check one) senile			Type of capsulotomy (check one)	_
<b>3</b> 7	(specify)		CCCR (continuous curvilinear capsulorhexis)	
Pathology (check yes or no for each)		t assessable	Other (specify)	
Pseudoexfoliation				
Glaucoma			Position of the loops (check one) in the bag	_
Previous glaucoma filtering surgery			partly in the bag	
Poor pupil dilation		1,40	in the sulcus	
Previous uveitis		iiC/F	uncertain	
Previous retinal detachment		$C_{\prime\prime}$		10
Diabetic retinopathy				 
Macular degeneration			If yes, specify:	_
Amblyopia		_	, 50, 5000	
Other (specify)	000		Problems during surgery (check yes or no for each) yes n	10
	Axial length	n mm	4	╗
-	<b>5</b>			╗
Target postoperative refraction			Posterior capsular opacity remaining	╗┃
Signed informed yes			<del>-</del>	╗┃
consent obtained:	DD MM	YY		ו⊏
			Other(specify)	⊐ I
6				
Operative report	ate of surgery		If investigation lens not implanted indicate reason:	
		MM YY		
Ophthalmic viscosurgical device ι	ısed yes □	no 🗆		
If yes, specify			Lens implanted. Place label here:	
Intraocular medication (check yes	or no for yes			
each)			Time incision to closure min.	
Adrenalin	_	_	Signature of investigator	$\dashv$
Acetylcholine			Signature of investigator	
Carbachol Other (apacity)				
Other (specify)	□		YY MM DD	
			1	

Table A.3 — Post-operative case report form for posterior chamber lens clinical investigation

Investigator name:					Clinical trial number	.r:	_
Patient number:	Patient initials:			D	ate of birth:		
Post-operative repo	rt				Other pathology and complications (Contin	nued)	
. oot operative rope	₹	MM DD				present	absent
Eye	rig	ht 🗆	left		Fibrin in pupil		
Check if the patient is una	available for this sched	luled examir	nation		Cortical remnants		
but continuing in the cl		gn form wi	th all		Nuclear remnants		
evaluation in form left blan	,				IOL optic decentration		
If the patient is discont primary reason:	tinued from the inves	stigation, in	dicate		if present: mm		
primary reason.					IOL optic tilt		
					if present:degrees		
Refraction	Sphere				IOL dislocation out of the posterior chamber		
	Cylinder				IOL optic discoloration		
	Axis				IOL optic opacities		
Keratometry	K1	_D D			Retinal detachment		
	K2	_			Diabetic retinopathy		
Boot corrected viewal co		Op. eye	Fello	W	Cystoid macular oedema		
Best corrected visual ac	uity		eye		if present diagnosed:		
					clinically		
or about one	Financian t		_		by fluorescein angiography   Macular degeneration	_	_
or check one	Finger count Hand movement				Optic atrophy		
	Light perception				Орис апорну	yes	no
	No light perception				Anterior capsular opacification present?		
IOP (application)	. to light perception	 mm Hg		W.	Is the posterior capsule intact?	_	_
Medications used up to t	this visit	topical	syste	emic	if intact:		
(check yes or no for each)		yes no	yes	no	posterior capsule fibrosis		
Corticosteroids					Elschnig's pearls		
Antibiotics					if not intact:		
NSAIDs					has the capsule been opened since		
Glaucoma medication	''م				the last reported visit?		
Other (specify)		<u> </u>			Other pathology?		
Corneal stromal oedema	all .	wound	cent	ral	specify:		
none	cO.						
mild/moderate	$\sim$ 0				If visual acuity less than 0,5 (20/40, 6/12	) indica	te main
severe			<u> </u>		reason:		
Iritis (check one)		none mild					· · · · · · · · · · · · · · · · · · ·
		moderate			Has the operated eye undergone any surgica	l ves	no
DE.		severe			reintervention since last reported visit?		
Other pathology and con (check present or absent for e		present	abse	ent	If yes, specify:		
(check present of absent for e	aon						
Wound leak					Has the national evangianced any adverse		20
Flat anterior chamber					Has the patient experienced any adverse event or ophthalmic adverse device effect?	yes 🗆	no
Hyphema							ш
Endophthalmitis							
if present	_					<b>.</b> .	
infectious					If yes, fill in the adverse event/ adverse devic form.	e effect i	report
sterile			П		ioni.		
Vitreous in anterior chamb Vitreous to wound	ei				If sorious, also contact the spensor in accorda	nco with	local
Raised IOP requiring treat	ment				If serious, also contact the sponsor in accorda regulations.	IIOC WILL	i iocai
Pupillary block	mont						
Anterior synechiae					Signature of investigator		
Posterior synechiae							
Deposits on IOL						TY MM	DD

# Table A.4 — Pre-operative/ operative case report form for anterior chamber lens clinical investigation

Investigator name:				Clinical tria	ıl number:	
Patient number: P	atient initials:		Sex: Male:	□ Race:	Caucasian	
			Female:		Black	
					Asian	
Date of birth:					Other	
YY MM DD					Mixed	
Pre-operative report			Irrigating solution used		yes	no
r re-operative report	TY	MM DD	irrigating solution used			
Operative eye	right □	left □	If yes, specify			
<u> </u>	Operative eye	Fellow eye	Periocular medication	yes (specify, if ap	nronriate)	no
Best corrected visual acuity	Operative cyc	1 CllOW Cyc	Anaesthetic			
or check one:			Antibiotic			) 🗆
Finger count			Corticosteroid			
Hand movement			Other (specify)		00	
Light perception			(0,000.,)		7.	_
No light perception			Incision		$\overline{\alpha}$	
IOP (applanation): On over	mmHa Follow	, ava:	Size mr	n 🔨		
IOP (applanation): Op. eye: _	IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	eye IIIIIIng	Type (e.g. corneal, limbal,	, scleral tunnel) 🤒	·	
Corneal status (check yes or no	for each)	yes no	Type of lens extraction (	check one)		
Normal	,		, ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,			
Guttata			Phacoemulsification	.00		
Other pathology (specify)			Other (specify)	داع		
Endothelial cell count (if done)	cells/mm <sup>2</sup>			0		
Corneal thickness (if measured	d): mm		Type of capsulotomy (ch	eck one)		
Cataract			d v			
	senile		CCCR (continuous curvilir	near capsulorhexis)		
	other (specify)	🗆	Other (specify)			
	,,		XO.			
Pathology (check yes or no for e	ach) yes no	not assessable			yes	no
Pseudoexfoliation			Other surgical procedur	es (check yes or no)		
Glaucoma			If yes, specify:			
Previous glaucoma filtering sui	rgery 🗆 🗆					
Poor pupil dilation			7.			
Previous uveitis		×O	Problems during surger	v (check ves or no for	each) yes	no
Previous retinal detachment		A.	Anterior segment bleeding		ĺ	
Diabetic retinopathy			Iris damage			
Macular degeneration			Posterior capsular opacity	remaining		
Amblyopia		_,	Posterior capsular rupture	1		
Other (specify)		$\omega$	Anterior vitrectomy	•		
Other (specify)	- ' ' <u> </u>	)`	Other (specify)			
Di	1/4 - 5	A - 2 = L L t l -				
Biometry	K1D	Axial length	If investigation lens not	implanted indicate	reason:	
Target postoperative refraction		mm				
			ļ <del></del>			
Signed informed consent	yes □	TY MM DD	Lens implanted. Place la	abel here:		
obtained:	<b>~</b>					
	Dete et comme		_			
Operative report	Date of surgery	/				
Implantation /						
primary			Lens orientation			
	ondary, specify reas	on	Lens orientation			
1 Secondary	maary, specify reas		Time to incision closure		minutes	
Ophthalmic viscosurgical de	vice used	yes □ no □			······································	
If yes, specify		,00				
Intraocular medication (check	voo or no for each	V00 00	Clausetone of located	larata u		
Adrenalin	yes or no lor each)	yes no □ □	Signature of investi	igator		
Acetylcholine						
Carbachol					TY MM DD	
Other (specify)						
\ 1			L			

Table A.5 — Post-operative case report form for anterior chamber lens clinical investigation

Investigator name	e:			Clinical trial numb	er:	_		
	Patient initials	S:		Date of birth:				
Post-operati	ve report	7 MM DD		Other pathology and complications (Continued)				
Eye	riç	ght 🗆 🗆	left □	Posterior synechiae Incorrect lens size	present	absent		
examination but form with all eval	ent is unavailable for this s continuing in the clinical ir luation in form left blank). iscontinued from the inves	nvestigation (s		Iris tuck Deposits on IOL Fibrin in pupil Cortical remnants Nuclear remnants IOL optic decentration if present: mm	<u> </u>			
Refraction	Sphere Cylinder		<del></del>	IOL tilt if present: degrees				
Keratometry	Axis K1 K2	 _D _D		IOL optic discoloration IOL optic opacities IOL dislocation out of the anterior chamber				
Best corrected or check one:		erative eye ——  □ □ □ □ □	Fellow eye  ——  □  □  □	Retinal detachment Diabetic retinopathy  Cystoid macular oedema if present diagnosed: clinically by fluorescein angiography				
IOP (applanation	1)	mm	n Hg	Macular degeneration				
Medication user (check yes or no for Corticosteroids Antibiotics NSAIDs Glaucoma medic	,	topical yes no		Is the posterior capsule intact? if intact:     posterior capsular fibrosis	yes	no -		
Other (specify) _		- Click		Elschnig's pearls  If not intact:				
Corneal stroma none mild/r sever	moderate	wound	central  □ □ □	has the capsule been opened since last reported visit? Other pathology? Specify:	_			
Iritis (check one	205150.	none mild moderate severe		If visual acuity less than 0,5 (20/40, 6/12) indice  Lens orientation	ate main r	reason:		
Other pathology (check present or a Wound leak	y and complications obsent for each)	present	absent	Has the operated eye undergone any surgical reintervention since last reported visit?  If yes, specify:	•	es no		
Flat anterior char Hyphema Vitreous in anter Vitreous to woun	ior chamber			Has the patient experienced any adverse event or ophthalmic adverse device effect?				
Hypopyon Iris atrophy Eversion of the p				If yes, fill in the adverse event/adverse device e  If serious, also contact the sponsor in accordance regulations.				
Endophthalmitis if present:	infectious □ sterile □			Signature of investigator				
Raised IOP requ Pupillary block Anterior synechia	iring treatment							

Table A.6 — Adverse event/ adverse device effect report form

Investigator name:		Clinical trial numbe	er:
Patient number:	Patient initials:	Date of birth:	
1. Operative eye	right □ left □	11. Treatment of adverse event (please print)	
2. Date of implant	TYY MM DD	11. Treatment of adverse event (please print)	3006
3. Lens model number	<del></del>	27.	
4. Lens serial number	<del></del>	. 2703	
5. Power of IOL	D	N	
Adverse event		12. Outcome (check one)	
6. Date of onset	DD MM YY	Complete recovery	
7. Duration (hours, days, et	c.)	Recovered with sequelee	
8. Severity of adverse ever	nt	Not recovered on day of report	
	(check one)	Died	
mild		Prognosis, if not recovered:	
moderate		:6 <sup>1</sup>	
severe		7,	
9. Describe course of adve	erse event (please print)	Comments (please print)	
	ROSIS	13. Does reporting physician believe adverse event is lens related?  Comments (please print)	yes no
10. Diagnosis of adverse e	vent (please print)		
		Signature of investigator Telephone	

# Annex B

(informative)

# **Evaluation of post-operative adverse event and visual acuity rates**

## **B.1 General**

In order to allow for an uncontrolled study, rates of adverse events and visual acuity were taken from data in USA studies to derive safety and performance endpoints (SPE).

# **B.2 Background**

The data for the SPE rates were derived from weighted averages of the data from large clinical investigations of anterior and posterior chamber IOLs.

The data for posterior chamber IOLs were taken from eight recent clinical investigations of posterior chamber IOLs that were approved in the US (December 1989 to December 1997). The pooled sample size for these clinical investigations was 4 210 for adverse events and overall best corrected visual acuity (BCVA), and 3 035 for best case BCVA.

The data for anterior chamber IOLs were taken from five recent clinical investigations for anterior chamber IOLs that were approved in the US (March 1988 to June 1991). The pooled sample size for these clinical investigations was 952 for adverse events and overall BCVA, and 635 for best case BCVA.

# B.3 Adverse event and visual acuity rates

The adverse event and visual acuity rates are provided in Tables B.1, B.2, B.3 and B.4.

For adverse events not included in this annex, comparison with published literature, previous clinical experience and the investigators clinical judgement, will determine acceptability.

Table B.1 — Anterior chamber IOL adverse event rates

	SPE	Numbe	r of subjects = 100	Number of subjects = 300		
Adverse event	rate <sup>c</sup>	Threshold rate <sup>d</sup>	Max. number of cases allowed before SPE rate exceeded <sup>e</sup>	Threshold rate <sup>d</sup>	Max. number of cases allowed before SPE rate exceeded <sup>e</sup>	
	%	%		%		
Cumulative:						
Cystoid macular oedema	10,0	18,8	15	14,9	39	
Hypopyon	0,2	3,0	1	1,4	2	
Endophthalmitis <sup>a</sup>	0,2	3,0	1	1,4	200	
Lens dislocated from anterior chamber	1,1	5,4	3	3,2	1.9	
Pupillary block	2,0	7,8	5	4,5	10	
Retinal detachment	1,2	5,4	3	3,4	7	
Secondary surgical intervention <sup>b</sup>	2,6	8,5	5	5,6	13	
Persistent:				0		
Corneal stroma oedema	0,5	4,2	2	2,2	4	
Cystoid macular oedema	3,8	10,1	7	7,1	17	
Iritis	0,9	5,4	3/4111	3,0	6	
Raised IOP requiring treatment	2,1	7,8	11/15	4,9	11	

<sup>&</sup>lt;sup>a</sup> Endophthalmitis is defined as inflammatory reaction (sterile or infectious) involving the vitreous body.

b Excludes posterior capsulotomies.

The SPE rate is the safety and performance endpoint.

The threshold rate is the minimum rate detectable as statistically significantly different from the SPE rate (greater than the SPE rate in the case of adverse events; less than the SPE rate in the case of BCVA).

The maximum number of cases allowed before SPE rate exceeded are the maximum number of subjects with that adverse event that can occur in a clinical investigation before the rate in that investigation becomes statistically significantly greater than the SPE rate.

Table B.2 — Posterior chamber IOL adverse event rates

	SPE	Numbe	r of subjects = 100	Number of subjects = 300		
Adverse event	rate <sup>c</sup>	Threshold rate <sup>d</sup>	Max. number of cases allowed before SPE rate exceeded <sup>e</sup>	Threshold rate <sup>d</sup>	Max. number of cases allowed before SPE rate exceeded <sup>e</sup>	
	%	%		%		
Cumulative:						
Cystoid macular oedema	3,0	8,9	6	6,0	14	
Hypopyon	0,3	3,0	1	1,8	3	
Endophthalmitis <sup>a</sup>	0,1	3,0	1	1,0	0001	
Lens dislocated from posterior chamber	0,1	3,0	1	1,0	1.72	
Pupillary block	0,1	3,0	1	1,0	1	
Retinal detachment	0,3	3,0	1	1,8	3	
Secondary surgical intervention <sup>b</sup>	0,8	4,2	2	2,6	5	
Persistent:						
Corneal stroma oedema	0,3	3,0	11 50 K O.	1,8	3	
Cystoid macular oedema	0,5	4,2	2	2,2	4	
Iritis	0,3	3,0	(N) 1	1,8	3	
Raised IOP requiring treatment	0,4	4,2	2	1,8	3	

Endophthalmitis is defined as inflammatory reaction (sterile of infectious) involving the vitreous body.

Table B.3 — Overall post-operative BCVA 0,5 (6/12, 20/40) or better

ORIC	SPE	Numbe	r of subjects = 100	Number of subjects = 300	
Cens type	rate <sup>a</sup> Threshold rate <sup>b</sup>		Min. number of cases allowed before less than SPE rate <sup>c</sup>	Threshold rate <sup>b</sup>	Min. number of cases allowed before less than SPE rate $^{\circ}$
9	%	%		%	
Anterior chamber IOL	80,4	69,6	74	74,3	230
Posterior chamber IOL	92,5	84,4	88	88,3	270

<sup>&</sup>lt;sup>a</sup> The SPE rate is the safety and performance endpoint.

b Excludes posterior capsulotomies.

The SPE rate is the safety and performance endpoints

The threshold rate is the minimum rate detectable as statistically significantly different from the SPE rate (greater than the SPE rate in the case of adverse events; less than the SPE rate in the case of BCVA).

The maximum number of cases allowed before SPE rate exceeded are the maximum number of subjects with that adverse event that can occur in a clinical investigation before the rate in that investigation becomes statistically significantly greater than the SPE rate.

b The threshold rate is the minimum rate detectable as statistically significantly different from the SPE rate (greater than the SPE rate in the case of adverse events; less than the SPE rate in the case of BCVA).

<sup>&</sup>lt;sup>c</sup> The minimum number of cases allowed before less than SPE rate are the minimum number of subjects with BCVA 0,5 or better that can occur in a clinical investigation before the rate in that investigation becomes statistically significantly less than the SPE rate.