The IRIS

ARTISAN® / ARTIFLEX®

- Optimal location for refractive correction
- The toughest tissue within the eye
  Phakic, Aphakic and Trauma
Iris structure

The mid-peripheral iris follows the same radial pattern as the vascular and nerve system of the eye. This pattern of tissues allows the clips of the Artisan/Artiflex to “weave” into the tissue. Photo: Dr Chazalon

Iris Anatomy

// Vascular supply
Arterial inflow & venous backflow - Radially oriented terminal end-arteries - No connection between individual end-arteries.

// Nerve system
Nerve system runs parallel to the vascular supply - No nerve / vascular damage caused by iris fixation of Artisan/Artiflex.

// Pigment layer
Blue eyes: deep brown pigment on rear iris surface only.

// Post removal Artisan After 6 yrs. in eye
No sign of pigment loss at rear side of the iris at the sites where Artisan was clipped.

Brown eyes: deep brown pigment on rear iris surface + pigment within the iris stroma.
Why the Iris?

Iris-fixated IOLs were developed as an alternative to reduce the occurrence of the problems that arose from angle fixated IOLs.

- The 1950s saw several designs: iris sphincter with anterior and posterior loops. These lenses led to progressive complications and use was abandoned, because they rested / were fixed on highly mobile lens sphincter.
- Dr. Jan Worst designed the “iris claw” concept in 1978 as an aphakic lens. Making mid-peripheral fixation ideal for placing an IOL.

NOTE: iris freckle does not move with pupil dilation; Making mid-peripheral fixation ideal for placing an IOL.

Iris fixated IOL history
not all “iris” IOLs are/were created equal

Iris-fixated IOLs were developed as an alternative to reduce the occurrence of the problems that arose from angle fixated IOLs.

- The 1950s saw several designs: iris sphincter with anterior and posterior loops. These lenses led to progressive complications and use was abandoned, because they rested / were fixed on highly mobile lens sphincter.
- Dr. Jan Worst designed the “iris claw” concept in 1978 as an aphakic lens. Making mid-peripheral fixation ideal for placing an IOL.

Artisan/Artiflex enduring technology
- Claw fixation method has not changed since the introduction.
- Artisan/Artiflex claws have a fine slot to capture or enclavate a small knuckle of mid-peripheral iris that is virtually immobile.
- Artisan/Artiflex optic is bridged over the mobile iris and pupil.
- Artisan/Artiflex will NOT rotate or tilt.
- Use of a small portion of iris for fixation has proven to create no clinical trauma.

"Iris Bridge" protects the endothelium
ARTISAN®/ARTIFLEX® - Enduring

Proven to be one of the world’s safest, most effective IOL designs, with the broadest applications of any IOL - Phakic, Secondary/Aphakic and Trauma

- The Artisan/Artiflex concept has the longest design history which is still in use.
- The Artisan/Artiflex corrects hyperopia, myopia, and astigmatism, and is used routinely for aphakic, secondary and trauma implantations.

Endothelial cell HEALTH with ARTISAN®/ARTIFLEX®

- Cataract surgery, and all corneal incisions (laser or knife), will damage/re-model endothelial cells (see matrix of peer review data).
- Manipulation of instruments, and IOLs, during cataract surgery, do not cause un-due concern with endothelial cell loss, and so it is with the Artisan lens - careful insertion is key to endothelial health.
- With age, the number of cells decreases at a rate of 0.6% per year after age 18. This means that after 10 years, a loss of approximately 6% could be found*.
- In a 10 year, peer reviewed** Artisan study, no endothelial cell loss of this magnitude was found. The data demonstrates there is no significant long-term corneal endothelial cell loss over time.
- No correlation was found between endothelial cell loss at 10 years and the preoperative anterior chamber depth, which supports the hypothesis that an anterior chamber depth of at least 3.0 mm is an adequate safety measure for the implantation of Artisan.

** Tahzib NG, Nuijts RM, Wu WY, Budo CJ. Long-term Study of Artisan Phakic Intraocular Lens Implantation for the Correction of Moderate to High Myopia; Ten-Year Follow-up Results. Ophthalmology 2007; 114(6):1133-42
# ARTISAN®/ARTIFLEX® vs. ICL & Standard Cataract

## Annual Endothelial cell loss

<table>
<thead>
<tr>
<th></th>
<th>ARTISAN/ARTIFLEX</th>
<th>ICL</th>
<th>Cataract surgery</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Annual Endothelial cell loss</strong></td>
<td>Only studies with &gt; 50 eyes of Myopia, Toric showed 1.18% loss(^1)</td>
<td>Only studies with &gt; 50 eyes of Myopia for ICL showed 1.19% loss(^1)</td>
<td>2.5% per year for at least 10 years after surgery, even without a lens implant(^3)</td>
</tr>
<tr>
<td><strong>Sizing</strong></td>
<td>One size fits all eyes exactly</td>
<td>4 sizes – fits no eye exactly</td>
<td>Sulcus issues</td>
</tr>
<tr>
<td><strong>Centration</strong></td>
<td>Surgeon’s choice</td>
<td>Anatomy decides</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Toric stability</strong></td>
<td>Does not rotate or tilt, after 24hr good vision(^2)</td>
<td>Can rotate and tilt and long visual recovery(^6)</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Incision size</strong></td>
<td>3.2 / 5.2 / 6.2 mm</td>
<td>3.2 mm</td>
<td>1.8 mm - 3.5 mm</td>
</tr>
<tr>
<td><strong>Clinical history</strong></td>
<td>25 yrs. fixation method has been unchanged</td>
<td>15 yrs. - design changed 5 times to address complications</td>
<td>NA</td>
</tr>
<tr>
<td><strong>Control of lens position</strong></td>
<td>Surgical learning curve, requires millimeters of clearance</td>
<td>Sizing, centration, limited clearance in sulcus - only microns of clearance in sulcus</td>
<td>Capsule/sulcus issues</td>
</tr>
<tr>
<td><strong>Main concern with design</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>


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# ARTISAN®/ARTIFLEX® vs. ICL - Diopter Ranges

**POWER RANGES (D)**

- **ARTISAN® Toric** | -22.0 to +14.0 D (Sphere) |
- **ARTISAN® Toric** | -22.5 to +10.0 D (Cylinder) |
- **ARTISAN® Myopia** | -23.5 to -1.0 D |
- **ARTISAN® Myopia** | -23.5 to -1.0 D |
- **ARTISAN® Aphakia** | +2.0 to +30.0 D |
- **ARTISAN® Hyperopia** | +10.0 to +12.0 D |
- **ARTISAN® Toric** | -7.5 to -1.0 D (Cylinder) |
- **ARTISAN® Toric** | -5.0 to -1.0 D (Cylinder) |
- **ARTISAN® Toric** | -13.5 to -1.0 D (Sphere) |
- **VISIAN ICL Toric** | +0.5 to +6.0 D (Cylinder) |
- **VISIAN ICL Toric** | +10.0 to +10.0 D (Cylinder) |
- **VISIAN ICL** | +10.0 to +10.0 D (Cylinder) |

Diopter ranges, according to 2016 publications
What is Quality of Vision?
All “20/20 vision” is not equal - Why?

A retrospective study* was performed comparing Optical Quality of Iris fixated Phakic IOL versus Sulcus Fixed Phakic IOL

Objective
With a validated objective tool, determine the differences in quality of vision (Optical Scatter Index (OSI)) for two different Phakic IOLs.

Conclusion
This study showed the iris fixated Phakic IOL produced the best quality of vision in all patients, and in some patients with sulcus fixated Phakic IOL the quality of vision was degraded equal to a +3 cataract.

* Pending publication; Dr. Lee, Korea

The visual acuity in these patients are nearly the same but the Quality of vision is very different. This objective data (OSI) shows the Quality of vision in the Artiflex is nearly 4 times better than in the ICL, even with visual acuities that are nearly identical.

Artiflex
VA = 1.24
OSI = .05

ICL
VA = 1.18
OSI = 2.3
Accommodation and Phakic IOLs

**ARTISAN / ARTIFLEX vs. ICL**

**ARTISAN / ARTIFLEX**
In a 3 year post-op multicenter peer review study the distance between the posterior surface of the Artisan/Artiflex and anterior surface of the crystalline lens was measured during accommodation. 1)

The study proved the distances remained constant with accommodation - this suggests that the iris diaphragm and crystalline lens act as a unit and move forward. 1)

ACD decreases with accommodation as a result of the forward movement of the iris diaphragm and crystalline lens. With the Artisan/Artiflex no measurement was found less than 2.0 mm at any point in the examination, which is considered the limit of safety for the corneal endothelium. 1)

**ARTISAN / ARTIFLEX allows natural accommodation to continue with age**

A main concern with all Phakic IOL is how they will interact with anterior segment structures (mainly anterior chamber angle, ciliary sulcus space, corneal endothelium, and crystalline lens). Modifications in the anterior segment can occur during accommodation and throughout life, and could predispose patients to premature cataract formation as they age, and/or limit the eyes ability to accommodate. With every diopter of accommodation the anterior pole of the crystalline lens moves forward 30µ. 1)

**ICL**
When we consider ICL sizing we usually think about width and not depth of sulcus. The ICL has very critical distance tolerances - consider the ICL's ability to block the natural lens ability to accommodate. The depth of the sulcus space shrinks as we age - an ICL that “fits” at 30 years old may not fit at 50 years old.

The ICL sizing criteria allows only microns of tolerance before problems can occur. The space allowance in the sulcus is very “tight”.

As the ICL will be located so close to the natural lens, some hesitation, with respect to the patient’s age would be wise. The natural lens grows with age and may contact the ICL in time. In young patients during accommodation the natural lens will move forward and risk to come into contact with the ICL. The ICL may even inhibit the patient’s ability to accommodate. Contact of the ICL with the natural lens may therefore cause early cataract formation.

1 José Luis Güell, MD, Merce Morral, MD, Oscar Gris, MD, Javier Gaytan, MD, Maite Sisquella, Opt, Felicidad Manero, MD

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Figure 1. Clinical photographs showing ARTISAN PIOL (A) and ARTIFLEX PIOL (B) positioned in the anterior segment. C and D: Visante OCT of the same patient with relaxed accommodation. Note the calipers (blue lines) used for anterior segment measurements. E and F: Visante OCT of the same patient with -3.0 D of accommodation. G and H: Visante OCT of the same patient with -7.0 D of accommodation.
**ARTISAN® Aphakia**

The ideal IOL for implantation in patients with aphakia without capsular support.

Based on the long term experience of iris fixation, the ARTISAN® Aphakia IOL is a predictable, safe, high precision implant.

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**Features & Benefits**

- Iris Fixation
- One size fits all
- Long term clinical experience
- Predictable, reliable, stable
- Also fit for retropupillary fixation
- Long term safety

---

**Specifications**

<table>
<thead>
<tr>
<th>ARTISAN® Aphakia IOL</th>
<th>Material</th>
<th>Total Ø</th>
<th>Body Ø</th>
<th>A-Constants*</th>
<th>AC Depth</th>
<th>Diopteric Powers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aphakic IOL Model 205</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>5.4 mm</td>
<td>Ultrasound: 115.0; Optical: 115.7 (SRK T)</td>
<td>3.3 mm</td>
<td>2.0 D to 30.0 D (1.0 increments)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Retropupillary: Ultrasound: 116.8; Optical: 116.9 (SRK T)</td>
<td></td>
<td>14.5 D to 24.5 D (0.5 increments)</td>
</tr>
</tbody>
</table>

*See www.ophtec.com for more IOL constants

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**Artisan Aphakic Benefits Matrix**

Aphakic, Trauma, Complications

<table>
<thead>
<tr>
<th></th>
<th>Artisan</th>
<th>Angle Supported lens</th>
<th>Sclera sutured PC IOL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Time</strong></td>
<td>10-20 minutes</td>
<td>10-20 minutes</td>
<td>20 to 60+ minutes</td>
</tr>
<tr>
<td><strong>Safety</strong></td>
<td>Complications limited to technique</td>
<td>Angle related complications</td>
<td>Sutures can erode and refraction unstable</td>
</tr>
<tr>
<td><strong>Outcomes</strong></td>
<td>Excellent, predictable</td>
<td>Angle related complications</td>
<td>Refraction not predictable, lens tilt, hemorrhage and secondary glaucoma</td>
</tr>
<tr>
<td><strong>Clinical History</strong></td>
<td>30+ years</td>
<td>Removed from many markets</td>
<td>30+ years</td>
</tr>
<tr>
<td><strong>Suturing of IOL required</strong></td>
<td>No</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td><strong>Surgical technique</strong></td>
<td>Easy</td>
<td>Easy</td>
<td>Complicated and extensive</td>
</tr>
<tr>
<td><strong>Fixation options</strong></td>
<td>Iris</td>
<td>Angle</td>
<td>Sclera, sulcus, iris</td>
</tr>
</tbody>
</table>

Artisan lenses are used in case of certain complications when PC lenses cannot be used:
- Insufficient support of the capsular bag
- Loss of capsular bag

**Trauma**

Secondary pathologies:
- Marfan’s Syndrome
- Pseudoexfoliation
- Congenital cataract
- Weil-Marchesani
- Homocystinuria
ARTISAN® Myopia and Hyperopia

ARTISAN® Myopia; first FDA approved Phakic IOL worldwide (2004)
ARTISAN® has passed the test of time by filling the need for those who seek a predictable and stable solution for the surgical correction of myopia, hyperopia and astigmatism.

Features & Benefits

- Iris Fixation
- Reversible treatment
- Predictable, reliable, stable, versatile
- Optimal clearance from vital tissues
- Various optical zone sizes
- Long term safety

// Specifications

<table>
<thead>
<tr>
<th>ARTISAN® PIOLs</th>
<th>Material</th>
<th>Total Ø</th>
<th>Body Ø</th>
<th>Dioptric Powers</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARTISAN® Myopia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.0 mm Model 206</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>5.0 mm</td>
<td>-1.0 D to -23.5 D (0.5 increments)</td>
</tr>
<tr>
<td>ARTISAN® Myopia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.0 mm Model 204</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>6.0 mm</td>
<td>-1.0 D to -15.5 D (0.5 increments)</td>
</tr>
<tr>
<td>ARTISAN® Hyperopia</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.0 mm Model 203</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>5.0 mm</td>
<td>+1.0 D to +12.0 D (0.5 increments)</td>
</tr>
</tbody>
</table>
ARTIFLEX® Myopia

Next Generation of Iris Fixated IOLs
ARTIFLEX® has a foldable lens body thus permitting a small incision. ARTIFLEX® offers a better predictability and faster recovery.

Features & Benefits

- Iris Fixation
- One size fits all
- Small incision, 3.2 mm; Controlled folding and unfolding
- Reversible treatment
- Aspherical edge design
- Optimal clearance from vital tissues
- Large optical zone

// Specifications

<table>
<thead>
<tr>
<th>ARTIFLEX® Myopia PIOL</th>
<th>Optic Material</th>
<th>Haptic Material</th>
<th>Total Ø</th>
<th>Body Ø</th>
<th>Dioptric Powers</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARTIFLEX® Myopia Model 401</td>
<td>Polysiloxane</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>6.0 mm</td>
<td>-2.0 D to -14.5 D (0.5 increments)</td>
</tr>
</tbody>
</table>
**ARTISAN® / ARTIFLEX® Toric**

**Only Artisan / Artiflex Toric doesn’t allow any rotation**

The ARTIFLEX® Toric PIOL is the latest extension of the successful ARTISAN® concept. It combines a spherical and cylindrical correction for low, moderate and high myopic eyes. The flexibility, extensive history and biocompatibility of the silicone optic enables implantation through a small incision, which results in almost no induced astigmatism as well as a fast recovery.

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**Features & Benefits: NO postop rotation**

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#### Specifications

<table>
<thead>
<tr>
<th>ARTISAN® Toric PIOL</th>
<th>Material</th>
<th>Total Ø</th>
<th>Body Ø</th>
<th>Dioptic Powers</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARTISAN Toric PIOL 0°</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>5.0 mm</td>
<td>Cylinder: -1 to -7.5 Sphere: -2.0 to +14.0 D (0.5 increments)</td>
</tr>
<tr>
<td>ARTISAN Toric PIOL 90°</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>5.0 mm</td>
<td>Cylinder: -1 to -7.5 Sphere: -2.0 to +14.0 D (0.5 increments)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ARTIFLEX® Toric PIOL</th>
<th>Optic Material</th>
<th>Haptic Material</th>
<th>Total Ø</th>
<th>Body Ø</th>
<th>Dioptic Powers</th>
</tr>
</thead>
<tbody>
<tr>
<td>ARTIFLEX Toric PIOL 0°</td>
<td>Polysiloxane</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>6.0 mm</td>
<td>Cylinder: -1.0 to -5.0 Sphere: -1.0 D to -13.5 D (0.5 increments)</td>
</tr>
<tr>
<td>ARTIFLEX Toric PIOL 90°</td>
<td>Polysiloxane</td>
<td>PMMA</td>
<td>8.5 mm</td>
<td>6.0 mm</td>
<td>Cylinder: -1.0 to -5.0 Sphere: -1.0 D to -13.5 D (0.5 increments)</td>
</tr>
</tbody>
</table>
ARTISAN® / ARTIFLEX® Instruments

// Artisan Instruments

OD 125  ARTISAN® / ARTIFLEX® Disposable Enclavation Needle

DO 240  ARTISAN® Reusable Enclavation Forceps

DO 270  ARTISAN® Reusable Implantation Forceps Refractive, Long
DO 272  ARTISAN® Reusable Implantation Forceps Refractive, Short

DO 274  ARTISAN® Reusable Implantation Forceps Cataract

DO 41  ARTISAN® Reusable Lens Manipulator Standard, straight

H 65.12.003  ArtiFix™ Reusable Holding Forceps

// Artiflex Instruments

OD 125  ARTISAN® / ARTIFLEX® Disposable Enclavation Needle

OD 110  ARTIFLEX® Disposable Insertion Spatula

OF 106  ARTIFLEX® Reusable Implantation Forceps Right

OF 105  ARTIFLEX® Reusable Implantation Forceps Left

OF 115  ARTIFLEX® Reusable Manipulator
**ARTISAN® / ARTIFLEX® Instruments VACUFIX™**

**Exact amount of tissue every time**

*Creates perfect reproducible “iris bridge”*

An enclavation system for the Artisan and Artiflex (Toric)(P)IOLs using the vacuum of your phaco machine to grasp a fold of iris tissue. The VacuFix consists of two disposable handles, one for the right side and one for the left. This will allow an optimal positioning and centration of the (T)(P)IOLs. The VacuFix tip with aspiration hole creates a perfect “iris bridge” with a controlled and reproducible amount of iris tissue. **Precision** for you and your patient is the key benefit, especially in toric Artisan/Artiflex surgery. For Artisan Aphakia cases the VacuFix adds **convenience**, as this system allows an easier grasp of iris tissue.

---

**// One System Fixates All**

- **Cataract Surgery**
  - ARTISAN® Aphakia IOL
- **Refractive Surgery**
  - ARTISAN® Myopia, Hyperopia & Toric PIOL;
  - ARTIFLEX® Myopia & Toric PIOL

**// Main Features & Benefits**

- **Vacuum enclavation for best positioning and centration of the (toric) (P)IOLs**
- **The VacuFix tip with aspiration hole creates a perfect “iris bridge”**
  - Fixed and reproducible amount of iris tissue
- **Preformed curved tip of the VacuFix makes it easy to reach the enclavation site**
- **The VacuFix is compatible with all phaco machines**
**SURGICAL PROCEDURE ARTISAN® PHAKIC IOLs**

*with VacuFix™ and Enclavation Needle*

1. Perform a main incision 0.1 mm larger than the optic diameter of the lens

2. Constrict the pupil; then introduce viscoelastic material, sodium hyaluronate (e.g. ArtiVisc or ArtiViscPlus)

3. Make paracenteses at 10 and 2 o’clock, pointing towards the fixation site. 1.2 mm (Enclavation Needle) 1.5 mm (VacuFix™) (In case of an Enclavation Forceps make two paracenteses 2.0-2.5 mm at 3 and 9 o’clock directed to the pupil)

4. Introduce the lens into the anterior chamber

5. Add some viscoelastic material on top of the lens

6. Rotate the lens into the horizontal position

7. Center the lens on the pupil; grasp the lens at the edge of the optic

8. Introduce the VacuFix through the paracentesis and make sure the hole of the VacuFix is placed underneath the slot of the claw. Create vacuum

9. Move the VacuFix with the occluded iris forward to the inferior part of the claw

10. Lift the VacuFix through the inferior claw and pull the VacuFix with the iris fold through the slot of the claw

11. Repeat the lens fixation to the iris on the other side

12. Make a “snow ploughing” movement by moving the Enclavation Needle downward and forward at the same time, creating a fold of iris tissue just under the claw of the lens

13. Make a peripheral iridotomy (or iridectomy), remove the viscoelastic material and close the main incision

*see www.ophtec.com for the Artisan Aphakic POSTERIOR CHAMBER (or RETROPUPILLARY) procedure.*
SURGICAL PROCEDURE ARTIFLEX®
with VacuFix™ and Enclavation Needle

1. Make paracenteses at 10 and 2 o’clock, pointing towards the fixation site.
2. Constrict the pupil; then introduce viscoelastic material, sodium hyaluronate (e.g. ArtiVisc or ArtiViscPlus).
3. Perform a main incision of 3.2 mm.
4-6 Attach the Artiflex PIOL to the Insertion Spatula.

7. Irrigate the lens with saline; introduce the lens into the anterior chamber with the Insertion Spatula.
8. Retract the Insertion Spatula; use a forceps to exert counter pressure.

9. Add some visco on top of the lens and rotate the lens into the horizontal position.
10. Center the lens on the pupil; grasp the lens at the superior claw with the Artiflex Holding Forceps.

11. Introduce the VacuFix through the paracentesis and make sure the hole of the VacuFix is placed underneath the slot of the claw; create vacuum.
12. Move the VacuFix with the occluded iris forward to the inferior part of the claw.

13. Lift the VacuFix through the inferior claw and pull the VacuFix with the iris fold through the slot of the claw.
14. Repeat the lens fixation to the iris on the other side.

14/15. Make a peripheral iridotomy (or iridectomy), remove the viscoelastic material and close the main incision.

11. Introduce the Enclavation Needle through the paracentesis.
12. Make a “snow ploughing” movement by moving the Enclavation Needle downward and forward at the same time, creating a fold of iris tissue just under the claw of the lens.
13. Repeat the lens fixation to the iris on the other side.

// VacuFix™

// Enclavation Needle
SURGICAL PROCEDURE ARTISAN® APHAKIA

Anterior fixation

1. Create a main incision which is 0.1 mm larger than the body diameter of the lens. The main incision should be at 90 degrees from the intended enclavation axis.

2. Constrict the pupil; then introduce viscoelastic sodium hyaluronate viscoelastics such as ArtiVisc® or ArtiVisc Plus®.

3. Make a paracentesis of 1.2 mm (Enclavation Needle) or 1.5 mm (VacuFix™) on each side of the main incision site.

4. Coat the IOL with viscoelastic prior to implantation to facilitate easy passing through the main incision.

5. Introduce the IOL into the anterior chamber through the main incision in a vertical orientation using the ARTISAN® Implantation forceps.

6. Rotate the lens 90 degrees and align with the enclavation axis using the ARTISAN® Lens Manipulator.

7. Bring the Artisan implantation forceps through the main incision, firmly grasp the IOL at the center of the optic and choose a side of the lens to enclavate. Center the IOL with the pupil.

8. Introduce the VacuFix™ through the paracentesis and make sure the hole of the VacuFix is placed underneath the slot of the claw. Create vacuum.

9. Move the VacuFix with the occluded iris forward to the inferior part of the claw.

10. Lift the VacuFix through the inferior claw and pull the VacuFix with the fold of iris tissue through the slot of the claw.

11. Repeat the lens fixation to the iris on the other side.

11/12. Make a peripheral iridotomy (or iridectomy), remove the viscoelastic material and close the main incision.

8. Introduce the Enclavation Needle through the paracentesis.

9. Make a “snow ploughing” movement by moving the Enclavation Needle downward and forward at the same time, creating a fold of iris tissue just under the claw of the lens.

10. Repeat the lens fixation to the iris on the other side.
1. Create a main incision which is 0.1 mm larger than the body diameter of the lens. The main incision should be at 90 degrees from the intended enclavation axis.

2. Inject a sufficient amount of high viscosity viscoelastic to maintain anterior and posterior chamber depth. Do not constrict the pupil.

3. Make a paracentesis of 1.2 mm on either side of the intended main incision site. The paracenteses must be in line with the haptics.

4. Invert (upside down) the IOL before insertion into the eye. Coat the IOL with viscoelastic prior to implantation to facilitate easy passing through the main incision.

5. Introduce the IOL into the anterior chamber through the main incision in a vertical orientation using the ARTISAN Implantation forceps.

6. Rotate the lens 90 degrees and align with the enclavation axis using the Artisan Lens Manipulator.

7. Bring the Artisan implantation forceps through the main incision, firmly grasp the IOL at the centre of the optic.

8. Carefully move the IOL through the non-constricted pupil.

9. Move the IOL upwards against the iris to facilitate enclavation and choose a side of the lens to enclavate.

10. Insert a cannula (for example 25 G) through one of the paracenteses, while securely holding the optic with the implantation forceps.

11. Sufficient iris tissue must be placed through the haptic slot to ensure adequate lens stability.

12. Repeat the previous steps at the other side of the lens. Make sure the lens is well centered and verify the amount of iris tissue which is enclavated.

13. Perform an iridotomy or iridectomy outside the periphery of the IOL. Remove all viscoelastic from the eye. Also flush out viscoelastic from underneath the IOL and close the main incision.
ARTISAN® / ARTIFLEX® Overview
Avoiding and Managing Complications;
Patient Selection Criteria - Proper Enclavation

All refractive procedures have a common concept: **Careful patient selection criteria are critical to successful outcomes.**

The ARTISAN/ARTIFLEX, like all refractive procedures, has “rules” to ensure success. We present here a consolidation of critical measurements. For a more comprehensive discussion please review the Ophtec Training Manual. Additionally, with refractive procedures, like with all surgical procedures, complications can happen. How to manage these is most important. We also address adverse events that could happen.

**ARTISAN**

*Endothelial health*

Minimum preoperative anterior chamber depth is 3.0 mm from epithelium, based on a minimum critical distance of 1.0 mm. (reference to critical distance tables in Ophtec Training Manual).

Minimum preoperative endothelial cell density (ECD) depending on patient age:

- < 25 years of age: 2800 cells/mm²;
- 26 to 30 years of age: 2650 cells/mm²;
- 31 to 35 years of age: 2400 cells/mm²;
- 36 to 45 years of age: 2200 cells/mm²;
- > 45 years of age: 2000 cells/mm²;

In order to assess the safety of the lens over time, patients should be examined 6 months after surgery and subsequently once a year. The follow-up examination should include monitoring of endothelial cell counts. Follow-up frequency should be increased to once every six months in case the decrease in cell count exceeds the physiological norm or when anterior chamber measurements show that the anterior chamber is becoming shallower due to developing cataracts. Patients should also be instructed not to rub the eye.

**Achieving good visual outcome**

- Accurate determination of preoperative refraction is crucial for achieving good refractive outcomes. A clinically significant difference between cycloplegic and manifest refraction is a contraindication.
- Pupil sizes in scotopic conditions should be ≤ 7.0 mm to reduce the risk of glare and halos.

**Avoiding complications**

An abnormally cone-shaped, bulging iris (typical for hyperopic eyes) is a contraindication, as it can cause formation of synechiae. For a complete overview of indications, contraindications and surgical technique see Ophtec Training Manual.

**ARTIFLEX**

*Endothelial health*

Minimum preoperative anterior chamber depth is 3.2 mm from epithelium, based on a minimum critical distance of 1.3 mm.

Minimum preoperative endothelial cell density (ECD) depending on patient age:

- < 25 years of age: 2800 cells/mm²;
- 26 to 30 years of age: 2650 cells/mm²;
- 31 to 35 years of age: 2400 cells/mm²;
- 36 to 45 years of age: 2200 cells/mm²;
- > 45 years of age: 2000 cells/mm²;

In order to assess the safety of the lens over time, patients should be examined 6 months after surgery and subsequently once a year. The follow-up examination should include monitoring of endothelial cell counts. Follow-up frequency should be increased to once every six months in case the decrease in cell count exceeds the physiological norm or when anterior chamber measurements show that the anterior chamber is becoming shallower due to developing cataracts. Patients should also be instructed not to rub the eye.

**Achieving good visual outcome**

- Accurate determination of preoperative refraction is crucial for achieving good refractive outcomes. A clinically significant difference between cycloplegic and manifest refraction is a contraindication.
- Pupil sizes in scotopic conditions should be ≤ 7.0 mm to reduce the risk of glare and halos.

**Avoiding complications**

- Select patient with flat irises. A convex, bulging or volcano shaped iris is a contraindication, as it can cause formation of deposits or synechiae.
- A correct surgical technique has to be used in order to avoid deformation of the PMMA haptics.

*If deposits on the optic surface appear*

Non-pigment deposits can be observed in a small percentage of patients – the exact reasons have not been established as this occurs rarely and follows no distinct pattern. The deposits are random and can appear in only one of the patient’s eyes and not in the fellow eye.

The deposits are usually observed between one and three months postop and slowly diminish after this period. There are no more deposits observed after one postoperative year, even in cases that were not treated with corticosteroids.
The discussions among the researchers suggest the probable cause of this phenomenon to be friction between the posterior lens surface and the iris, along with an implantation technique that is more difficult than that of the ARTISAN lens. However we have not been able to confirm the root cause due to very limited occurrence and randomness.

The following conclusions/guidelines were elicited from these discussions:

1. Eyes with a shallow ACD (<3.2mm) should be avoided. It is additionally of great importance that the iris is flat. Eyes with a convex, bulging or volcano shape must definitely be avoided. It is also advised that the pre-op examination techniques such as Scheimpflug photography or a OCT scan should be used. Unfortunately, it seems that not every suitable ARTISAN candidate is automatically a suitable ARTIFLEX candidate as well. Also, the iris of patients with myopia is not always evenly flat.

2. Excessive manipulation during surgery can lead to more deposits, but taking the learning curve of the technique into account, this should improve after a number of implantations. Enclavating an iris fold that is too large in the claw should be avoided. A large fold causes the lens to adhere more tightly to the iris. Use of the VacuFix will ensure the exact amount of iris tissue is enclavated every time.

3. Administration of preventative corticosteroids should be started after the implantation. This treatment must be maintained for four weeks. A schedule is cited here below. Some doctors administer a depo injection at the end of the implantation procedure.

**ARTIFLEX postoperative medication:**

**Antibiotics:**
1 drop of topical antibiotics 3 times daily during the first postoperative week, gradually reduced during 2 weeks.

**Corticosteroids:**
1 drop of strong-working topical steroids (for example, dexamethasone or fluormetholon) 3 times daily during the first four postoperative weeks. A peroperative depo injection of Depo-Medrol is optional.

**To prevent excessive eye pressure:**
Diamox or Betagan as needed.

In the event that a patient does develop deposits that impede vision, the doctor can treat the deposits with a brief course of corticosteroid treatment. This should cause the deposits to quickly disappear and the vision to recover.

If the deposits do return or do not disappear, a re-enclavation should be considered. In some cases, this has been known to stop these returning symptoms. If the symptoms do not stop and continue to return, this can lead to an explantation.

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**How to properly Enclavate the iris**

**PROPER technique:** Enclave iris tissue by bringing the iris tissue through the opposite “claw” from forceps

**INCORRECT technique:** DO NOT enclave the iris where the forceps is being held

See damage caused by incorrect enclavation
Surgeon Testimonials

**Dr I. Ahmed**, Canada

“The iris claw permits stable fixation of the lens, performed in a very efficient manner with minimal tissue manipulation or suture requirement”

**Dr Chul Myong Choe**, South Korea

“I have implanted over 10,000 ArtiLenses PIOLs and keep on going. The satisfaction comes from my patients. Nearly all of my patients thank me for giving them a new life”

**Dr S. Fukuoka**, Japan

“The ARTISAN® and ARTIFLEX® lenses are especially reliable to use in cases with astigmatism because the IOLs can be fixated stable on the iris without rotation”

**Dr V. Ghanem**, Brazil

“I decided to have Artisan lenses implanted in my own eyes. Today having experienced the patient position makes me feel even more confident about using Artisan lenses.”

**Dr J.L. Güell**, Spain

“Our 5 year experience with the ARTIFLEX® has been extremely positive, being the earlier refractive rehabilitation the main advantage over the rigid model. Hopefully, the long term (10/15 year) safety data will be as good as the one that we have had with the PMMA implant”

**Dr Huang Wei Jen**, Taiwan

“Once I trusted that the results were very good, I jumped at the opportunity of using ARTISAN® and, about a year ago, ARTIFLEX®”

**Dr L. Izquierdo**, Peru

“The Artisan/Artiflex lens is the most proven Phakic IOL platform in the world that I trust to use for my patient”

**Prof. Cheon-Ki Joo**, South Korea

“I can observe the whole aperture of the lens in the patient’s eye during the follow up period. So, I understand a patient’s condition completely. I think it would be the main advantage of the iris claw phakic IOL compared with a posterior chamber phakic IOL”

**Dr M. Landesz**, The Netherlands

“The ARTIFLEX® has the same ‘wow-effect’ one day postoperative as the lasik has”

**Dr R. Moreno**, Chile

“The Artiflex Lens has very good refractive predictability and rapid visual recovery. Its rotational stability make it very safe for toric correction. Also, there is no problem with the sizing of its length”

**Dr R. Ruiz Mesa**, Spain

“With the ARTIFLEX® Toric Lens and its option to be implanted through only 3 mm, at last I’ve found a phakic lens, which I have long needed for my patients: guaranteeing ease of procedure, rapid visual recovery and especially, safety of rotational stability independent of the calculation”

**Dr R. Spirig**, Switzerland

“ARTIFLEX® implantation is absolutely astigmatism neutral. The ARTIFLEX® can therefore be implanted not only in cases with high myopia, but also in cases with low myopia, where it is especially important to avoid any surgery-induced astigmatism. We experienced a very high degree of satisfaction in this group of patients who prefer to have refractive surgery with a reversible procedure rather than with a Lasik method”

**Dr Prof. M. Tetz**, Germany

“Due to my many years of experience in the pathology of intraocular lenses, I consider the iris fixation as a minimal traumatic fixation option in the phakic eye. With proper fixation, “sizing the IOL” is never a problem, unlike chamber-angle or posterior chamber intraocular lenses. Visual rehabilitation can be achieved very quickly and safely with the ARTIFLEX® IOL”
Surgical Videos

Use QR code or visit www.youtube.com/ophtecbv

ARTISAN Aphakia, the ideal IOL for secondary implantation
By Dr. J. L. Güell

ARTISAN PIOL implantation technique
By Dr. C. Budo

ARTIFLEX implantation; Lateral
By Dr. J. L. Güell

ARTISAN toric implantation; Step by Step
By Dr. J. L. Güell

ARTIFLEX toric implantation; Step by Step
By Dr. J. L. Güell

Needle enclavation of the ARTIFLEX PIOL

ARTIFLEX Toric with VacuFix
By Dr. J. L. Güell
Official ARTISAN & ARTIFLEX training courses

To obtain the best results with Artisan and Artiflex intraocular lenses, OPHTEC continuously conducts specialized ArtiLens certification courses and wetlabs throughout the world. By providing lenses only to certified professionals, OPHTEC guarantees the constant high quality of the results with the product.

Please check www.ophtec.com for training courses or send an e-mail to training@ophtec.com.