ENHANCING THE SAFETY PROFILE OF CATARACT PROCEDURES WITH THE ACTIVE SENTRY HANDPIECE AND INTREPID HYBRID TIP

Help limit complications with a combination of sensitive, automated IOP control and polymer technology phaco tip.

BY DAVID LUBECK, MD

Over the last 2 decades, we have been gradually reinventing phacoemulsification. The first technological change that had a significant impact on my approach was torsional ultrasound with the INFINITI Vision System (Alcon), which really helped me rethink my procedure. Next, the CENTURION Vision System (Alcon) Active Fluidics allowed me to select and maintain IOP so I could define and plan precisely what I wanted to do in each procedure and carry out the plan safely and efficiently. Now the CENTURION Vision System with ACTIVE SENTRY Handpiece combines ACTIVE FLUIDICS (Alcon) technology with a pressure sensor in the handpiece that measures intraoperative IOP directly. Also available is a polymer-ended phaco tip, the INTREPID Hybrid Tip (Alcon), which is designed to reduce the risk of capsular tears. The combination has made my phacoemulsification procedure safer, plus more efficient and controlled than ever before.

AUTOMATED FINE CONTROL OF IOP

Before the CENTURION, operating at or near physiologic IOP was not safely feasible. When we got the CENTURION, the ability to operate at physiologic IOP was novel at first, but I found the transition to be very intuitive. It was certainly compelling because we are always trying to achieve normal visual physiological optics and normal physiologic pressure management in all disease states. Remaining at physiologic levels during surgery supports those goals.

With the introduction of the ACTIVE SENTRY Handpiece (Alcon), not only was it feasible to operate at or near physiologic IOP, but it also became possible to refine IOP to precise levels tailored to different patients. With ACTIVE SENTRY, continuous IOP monitoring, leakage compensation, and automated patient eye level (PEL) provide a noticeably more stable intraocular environment and mitigate instability that can arise. The handpiece gives me an even greater degree of subtlety, precision, and scalability. It allows me to approach any lens with a great deal of forethought and planning, thus helping ensure my confidence in the success of the procedure.

By offering precise control, the ACTIVE SENTRY Handpiece improves the safety profile of phacoemulsification. That new base line in safety advantage is enhanced tremendously with another remarkable innovation, the INTREPID Hybrid Tip.

THE ADDED SAFETY OF POLYMER TECHNOLOGY

Designed to lower the risk of damage to the capsular bag and other tissues, the INTREPID Hybrid Tip has a polymer end with no sharp or metal edges. The ultrasonic and fluidics performance are similar to the all-metal INTREPID Balanced Tip. Because of the decreased risk of capsular rupture with the Hybrid Tip, phacoemulsification can be performed closer to the capsule and/or with higher flow and vacuum levels. This can help surgeons operate more comfortably with their existing techniques and give them the confidence to evolve their procedures.

We always welcome ways to reduce the rate of capsule rupture to help in safer procedures with better outcomes. Used together, the ACTIVE SENTRY Handpiece and INTREPID Hybrid Tip are effective innovations I’ve seen in my career to decrease the incidence of capsule rupture.
I also use the INTREPID Transformer I/A Handpiece (Alcon), which has the same polymer technology. Using polymer-tipped instruments throughout the procedure gives a sense of ease to all steps of lens removal. I can focus more closely on the subtleties or complexities of procedures instead of focusing on the elements of risk. For example, during phacoemulsification, I’m less frequently trying to assess how close the instrument is to the capsule or iris. By changing my focus, I can look at broader aspects of the procedure, such as nucleus position, capsular stability, and fluidic dynamics. The all-polymer procedure feels like the wave of the future. It changes the way I interpret the procedure moment to moment, making it a richer, more fulfilling experience and a more efficient procedure overall.

Because the INTREPID Hybrid Tip enhances my ability to more confidently disassemble the nucleus with an increased safety profile, I choose to use it in all but the very densest of nuclei (up to grade 4). By marrying the ACTIVE SENTRY Handpiece and INTREPID Hybrid Tip, I get the greatest possible flexibility and versatility from the CENTURION Vision System platform.

CASE: SAFER, MORE EFFICIENT CATARACT SURGERY

I performed cataract surgery on a 77-year-old woman with 2 to 3+ nuclear sclerosis and 2+ cortical changes.

A tapered sideport incision was made with a 1-mm blade. A square 2.2-mm clear corneal incision was made temporally. DisCoVisc OVD (Alcon) was used throughout the procedure. The approximately 5-mm capsulorhexis was centered nasally 0.5 mm. Cortical hydrodissection was performed until the nucleus could spin freely within the capsular bag.

For phacoemulsification, I used the INTREPID Hybrid Tip, which has the familiar Balanced Tip design with an innovative polymer end. There are no metal edges on this tip, so at no point can metal contact the capsule or other intraocular structures (Figure 1). This makes the tip much less likely to cause damage to the capsule or iris. The performance of this tip is similar to its all-metal counterpart; it grooves very efficiently for divide and conquer, and it holds the nucleus well for a chopping technique.

► ACTIVE SENTRY adjustments: Throughout the entire procedure, a sensor in the ACTIVE SENTRY Handpiece directly and continuously monitored IOP and managed the target IOP of 36. This technology also automates the patient eye-level function so that I do not need to sit the patient at any particular height to achieve the desired IOP for surgery. The handpiece also adjusts for average incision leakage, making the chamber significantly more stable. This feature has allowed me to raise my aspiration and vacuum levels, permitting a more efficient and predictable nucleus disassembly and removal. In this case, ACTIVE SENTRY engaged 11 times during phacoemulsification. This means that, at 11 different moments, the technology prevented or mitigated a surge event.

► Cortex removal and polishing: Next, to remove the cortex, I used the INTREPID Transformer I/A Handpiece, which is also
a novel and commendable innovation. In its primary mode, it is a coaxial polymer-tipped I/A handpiece. The polymer tip design allows me to place the tip up against the anterior and posterior capsules. When access to the subincisional cortex is required, the handpiece separates, and I insert the aspiration tip through the 1-mm sideport incision.

In this bimanual mode, I can easily assess the complete circumference of the capsular bag and polish the entire anterior capsule. If the capsule is at all unstable, then the bimanual mode facilitates easier cortex removal. When posterior capsular polishing is needed, the capsule can be drawn into the aspiration port and the tip passed across the capsular plane, yielding the clearest possible capsule.

Finally, I inserted the new lens. The viscoelastic was aspirated from the anterior chamber and behind the lens, finishing the cataract surgery.

**CASE: A SMOOTH PROCESS WITH AN INTRAOPERATIVE ADJUSTMENT**

Using the LENSX femtosecond laser (Alcon) I performed surgery on a 72-year-old woman with a 2 to 3+ nuclear and 2+ cortical cataract.

After opening the incisions, the capsule was drawn centrally to make sure that it was free and had no microadhesions. Hydrodissection was performed gently and slowly until all gas had been released anteriorly and the nucleus could be rotated within the capsular bag. Phacoemulsification was performed using the ACTIVE SENTRY Handpiece for fluidics management and the INTREPID Hybrid Tip, a balance-designed tip with a polymer end. The polymer tip’s efficiency is similar to a metal balanced tip, so it engages the nucleus and holds easily for division and emulsification of individual segments.

**Maintaining target IOP with ACTIVE SENTRY:** The ACTIVE SENTRY IOP Handpiece sensor continually, directly monitors IOP throughout the procedure, communicating with ACTIVE FLUIDICS. The system automates patient eye level, so it’s no longer necessary to set that on the panel or position the patient at a certain height. It also adjusts for average leakage from incisions. All of this occurs automatically in the background, engaging intermittently as needed to mitigate surge events and maintain chamber stability.

In this case, where the lens is graded a 2 to 3+, it was emulsified very efficiently. ACTIVE SENTRY allowed me to increase the flow and vacuum rates by as much as 30% and maintain a very stable chamber, even working at a near-physiologic IOP of 36 mm Hg (Figure 3).

Phacoemulsification required a CDE of 5.89. The ACTIVE SENTRY engaged a total of seven times during nucleus removal.

**Cortex removal and polishing:** Irrigation and aspiration of the cortex were done with the INTREPID Transformer I/A Handpiece, another Alcon polymer-tipped aspiration instrument (Figure 4). This handpiece starts as a coaxial I/A and converts to a bimanual if needed. The second handpiece is disengaged from the back of the first, and the aspiration tip is inserted through the 1-mm sideport incision. It can be used to polish the posterior capsule and remove any cortex that was inaccessible with the handpiece in coaxial mode. The aspiration port on the transformer bimanual handpiece has an oval contour that’s ideal for polishing the capsule. For the posterior capsule, polishing can be achieved by gently engaging the capsule with the tip and then dragging the tip slowly across the capsular surface.

**Intraoperative aberrometry and lens implantation:** The anterior chamber and capsular bag were filled with ProVisc OVD (Alcon)
and the pressure raised to the appropriate level. I turned off the microscope light, and the patient fixated on the red light within the ORA SYSTEM (Alcon) so I could take several series of measurements. A lens was chosen and implanted to provide the desired outcome.

Once again, I used the INTREPID Transformer I/A Handpiece for aspiration of the viscoelastic. The polymer tip is ideally designed to go behind the lens and safely work close to the capsule. It’s also an excellent lens manipulator, so it can be used to center and rotate the IOL.

The combination of the ACTIVE SENTRY Handpiece and ACTIVE FLUIDICS, with the polymer technology of the INTREPID Hybrid Tip and INTREPID Transformer I/A Handpiece, results in a very efficient and improved procedure. With precise control of IOP throughout surgery and lower risk of injury from the polymer-tipped instruments, I can focus on surgery with less concern about inducing complications.

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**CENTURION® Vision System Important Product Information**

Federal (USA) law restricts this device to sale by, or on the order of, a physician.

As part of a properly maintained surgical environment, it is recommended that a backup IOL injector be made available in the event the AutoSert® IOL Injector Handpiece does not perform as expected.

**INDICATION:** The CENTURION® Vision system is indicated for emulsification, separation, irrigation, and aspiration of cataracts, residual cortical material and lens epithelial cells, vitreous aspiration and cutting associated with anterior vitrectomy, bipolar coagulation, and intraocular lens injection. The AutoSert® IOL Injector Handpiece is intended to deliver qualified AcrySof® intraocular lenses into the eye following cataract removal.

The AutoSert® IOL Injector Handpiece achieves the functionality of injection of intraocular lenses. The AutoSert® IOL Injector Handpiece is indicated for use with the AcrySof® lenses SN60WF, SN6AD1, SN6AT3 through SN6AT9, as well as approved AcrySof® lenses that are specifically indicated for use with this inserter, as indicated in the approved labeling of those lenses.

**WARNINGS:** Appropriate use of CENTURION® Vision System parameters and accessories is important for successful procedures. Use of low vacuum limits, low flow rates, low bottle heights, high power settings, extended power usage, power usage during occlusion conditions (beeping tones), failure to sufficiently aspirate viscoelastic prior to using power, excessively tight incisions, and combinations of the above actions may result in significant temperature increases at incision site and inside the eye, and lead to severe thermal eye tissue damage.

Good clinical practice dictates the testing for adequate irrigation and aspiration flow prior to entering the eye. Ensure that tubings are not occluded or pinched during any phase of operation.

**AES/COMPLICATIONS:** Inadvertent actuation of Prime or Tune while a handpiece is in the eye can create a hazardous condition that may result in patient injury. During any ultrasonic procedure, metal particles may result from inadvertent touching of the ultrasonic tip with a second instrument. Another potential source of metal particles resulting from any ultrasonic handpiece may be the result of ultrasonic energy causing micro abrasion of the ultrasonic tip.

**ATTENTION:** Refer to the Directions for Use for the accessories/consumables and Operator’s Manual for a complete listing of indications, warnings, cautions and notes.

**Important Product Information for DisCoVisc® OVD**

**CAUTION:** Federal (USA) law restricts this device to sale by or on the order of a physician.

**INDICATIONS:** DisCoVisc® (Sodium Chondroitin Sulfate-Sodium Hyaluronate) Ophthalmic Visco-surgical Device (OVD) is indicated for use during surgery in the anterior segment of the eye. It is designed to create and maintain space, to protect the corneal endothelium and other intraocular tissues and to manipulate tissues during surgery. It may also be used to coat intraocular lenses and instruments during cataract extraction and IOL insertion.

**WARNINGS/PRECAUTIONS:** Failure to follow assembly instructions or use of an alternate cannula may result in cannula detachment and potential patient injury. Precautions are limited to those normally associated with the surgical procedure being performed. Although sodium hyaluronate and sodium chondroitin sulfate are highly purified biological polymers, the physician should be aware of the potential allergic risks inherent in the use of any biological material.

**ADVERSE REACTIONS:** DisCoVisc® Ophthalmic Viscosurgical Device was very well tolerated in nonclinical and clinical studies. A transient rise in intraocular pressure in the early postoperative period may be expected due to the presence of sodium hyaluronate, which has been shown to affect such a rise. It is therefore recommended that DisCoVisc® OVD be removed from the anterior chamber by thorough irrigation and/or aspiration at the end of surgery to minimize postoperative IOP increases.

**ATTENTION:** Reference the Directions for Use for a complete listing of warnings and precautions.

**Important Product Information for PROVISC® OVD**

**CAUTION:** Federal (USA) law restricts this device to sale by, or on the order of, a physician.

**INDICATION:** PROVISC® (Sodium Hyaluronate) Ophthalmic Viscoelastic Device (OVD) is indicated for use as an ophthalmic surgical aid in the anterior segment during cataract extraction and intraocular lens (IOL) implantation. Ophthalmic viscoelastics serve to maintain a deep anterior chamber during anterior segment surgery allowing reduced trauma to the corneal endothelium and surrounding ocular tissues. They help push back the vitreous face and prevent formation of a flat chamber during surgery.

**WARNINGS/PRECAUTIONS:**

- Postoperative increases in intraocular pressure have been reported with sodium hyaluronate products. The IOP should be carefully monitored and appropriate therapy instituted if significant increases should occur. It is recommended that PROVISC® OVD be removed by irrigation and/or aspiration at the close of surgery. Do not overfill anterior chamber. Although sodium hyaluronate is a highly purified biological polymer the physician should be aware of the potential allergic risks inherent in the use of any biological material; care should be used in patients with hypersensitivity to any components in this material. Cannula assembly instructions should be followed to prevent patient injury.

**ADVERSE EVENTS:**

- Postoperative inflammatory reactions such as hypopyon and iritis have been reported with the use of ophthalmic viscoelastics, as well as incidents of corneal edema, corneal decompensation, and a transient rise intraocular pressure.

**ATTENTION:** Reference the directions for use for a complete listing of indications, warnings and precautions.

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