Adding CONTOURA Vision to the Refractive Armamentarium

The following article highlights key questions addressed during a roundtable discussion of the role of personalized topography-guided LASIK among today's selection of refractive treatments. Led by Kerry D. Solomon, MD, the panel included Ronald R. Krueger, MD; Karl G. Stonecipher, MD; and R. Doyle Stulting, MD, PhD. Video of the roundtable can be viewed at eyetube.net/collections/contoura-pearls.

What are some advantages of CONTOURA Vision topography-guided treatments versus wavefront-optimized or wavefront-customized treatments?

Ronald R. Krueger, MD: I think there are two primary advantages. First, topography-guided treatments offer us an opportunity to eliminate most aberrations on the cornea. Because the cornea is the eye's first refractive structure, it is essential that we do everything we can to ensure that the front surface has the most uniform prolate shape possible, able to bring light to a single focus without bending. In addition to offering patients immediate visual benefits, a clear cornea that is free of aberrations is likely to have good long-term health and clarity until the patient requires lens replacement in the future.

The second advantage of topography-guided LASIK is that it does not induce aberrations. Wavefront-guided treatments allow us to measure and treat aberrations. They reduce the overall induced aberrations compared with conventional LASIK, but they still induce aberrations. Patients are never aberration-free. A myopic patient who has wavefront-optimized ablation, for example, is likely to have nearly the same visual outcomes we would expect from CONTOURA Vision, but without the same reduction of higher-order and even lower-order aberrations seen with CONTOURA Vision.¹

With topography-guided LASIK, there is a whole new outlook on postoperative aberrations. While most of us focus on visual outcomes and not higher-order aberration reduction, we do see improvements in symptoms and best corrected vision. That tells us the CONTOURA Vision procedure is going beyond the visual outcomes we accomplish with wavefront-optimized and wavefront-guided techniques.

Dr. Stulting, you were the medical monitor of the CONTOURA Vision FDA study. How did your research affect your view of this modality?

R. Doyle Stulting, MD, PhD: The study evaluated the efficacy of CONTOURA Vision for myopia and myopic astigmatism.² It was limited to “normal eyes,” excluding any subjects who had asymmetrical bowties, skewed radial axes, or any other features that could be considered abnormal. When we were planning the study, I was sure that the outcomes for these “normal eyes” would be no different than those we were accustomed to seeing with existing, approved treatments.

To my surprise—and that of everyone involved—the outcomes were much better than we could expect from standard treatments. The visual acuities were superb. For 30% of patients, postoperative uncorrected distance visual acuity was actually better than preoperative best spectacle-corrected distance visual acuity. Furthermore, subjects noted less light sensitivity, difficulty driving at night, reading difficulty, glare, halos, and starbursts than they did before surgery. These findings allow us to tell our patients that CONTOURA Vision may provide them with better uncorrected vision than they have with their glasses.

These exceptional outcomes can probably be attributed to CONTOURA Vision’s lack of dependence on pupil size/location, its ability to measure peripheral corneal aberrations that may be significant in low-light situations, its reproducibility, and its...
Adding CONTOURA Vision to the Refractive Armamentarium

What will your selection of LASIK treatment options look like going forward?

Kerry D. Solomon, MD: I am very accustomed to wavefront-guided technology, and most of the surgeons I know are entrenched in the mindset that wavefront-guided, wavefront-optimized, or wavefront-customized treatment is best. My one pearl for all of us is to question the status quo. Look at the data for CONTOURA Vision. See if something new and different makes sense in this case. Compared to wavefront procedures, CONTOURA Vision induces fewer aberrations and gives patients a clear cornea with less tissue ablation and possibly better contrast sensitivity. It certainly makes sense to me to try a procedure with those outcomes, and I think that a lot of other wavefront surgeons out there would reach the same conclusion.

Dr. Krueger: Ultimately, once we have the CONTOURA Vision technology, we can explain to patients that our practice has a full menu of different customized options, which allows us to choose the best procedure to achieve the results they want. Testing steers us in the right direction. If we obtain good quality topographic (Topolyzer Vario; Alcon) maps, we will perform CONTOURA Vision for the best results. If the topographic maps are not good, we might choose wavefront-optimized LASIK or perform wavefront-guided treatment to address some higher-order aberrations. The patient’s goals and expectations, combined with testing, drive individualized choices drawn from the full menu of options.

Karl G. Stonecipher, MD: To obtain the best possible visual outcomes for all of our patients, we have to design the most effective optical system for each one. That is the great thing about this robust surgical armamentarium. I will still want to do wavefront-optimized treatment for some patients, as well as perform wavefront-guided LASIK occasionally when I think it best. By offering another strong tool in this armamentarium, CONTOURA Vision makes customized treatment that much easier.

WaveLight® Eximer Laser System Important Product Information

This information pertains to all WaveLight® Eximer Laser Systems, including the WaveLight® ALLEGRETTO WAVE®, the ALLEGRETTO WAVE® Eye-Q, and the WaveLight® EX500.

Caution: Federal (U.S.) law restricts the WaveLight® Excimer Laser Systems to sale by or on the order of a physician. Only practitioners who are experienced in the medical management and surgical treatment of the cornea, who have been trained in laser refractive surgery (including laser calibration and operation) should use a WaveLight® Eximer Laser System.

Indications: FDA has approved the WaveLight® Eximer Laser Systems for use in laser-assisted in situ keratomileusis (LASIK) treatments for:

- the reduction or elimination of myopia of up to -12.00 D and up to 6.00 D of astigmatism at the spectacle plane;
- the reduction or elimination of hyperopia up to +6.00 D with and without astigmatic refractive errors up to 5.00 D at the spectacle plane, with a maximum manifest refraction spherical equivalent of +6.00 D;
- the reduction or elimination of naturally occurring mixed astigmatism of up to 6.00 D at the spectacle plane; and
- the wavefront-guided reduction or elimination of myopia of up to -7.00 D and up to 3.00 D of astigmatism at the spectacle plane.

In addition, FDA has approved the WaveLight® ALLEGRETTO WAVE® Eye-Q Eximer Laser System, when used with the WaveLight® ALLEGRETTO Topolyzer® and topography-guided treatment planning software for topography-guided LASIK treatments for the reduction or elimination of up to -9.00 D of myopia, or for the reduction or elimination of myopia with astigmatism, with up to -8.00 D of myopia and up to 3.00 D of astigmatism. The WaveLight® Eximer Laser Systems are only indicated for use in patients who are 18 years of age or older (21 years of age or older for mixed astigmatism) with documentation of a stable manifest refraction defined as ≤0.50 D of preoperative spherical equivalent shift over one year prior to surgery, exclusive of changes due to unmasking latent hyperopia.

Contraindications: The WaveLight® Eximer Laser Systems are contraindicated for use with patients who:

- are pregnant or nursing;
- have a diagnosed collagen vascular, autoimmune or immunodeficiency disease;
- have been diagnosed keratoconus or if there are any clinical pictures suggestive of keratoconus;
- are taking isotretinoin (Accutane®) and/or amiodarone hydrochloride (Cordarone®);
- have severe dry eye;
- have corneas too thin for LASIK;
- have recurrent corneal erosion;
- have advanced glaucoma; or
- have uncontrolled diabetes.

Warnings: The WaveLight® Eximer Laser Systems are not recommended for use with patients independence from lenticular aberrations. I believe we are seeing the benefits of optical correction of topographical abnormalities that are subtler than those we typically recognize.

Adding CONTOURA Vision to the Refractive Armamentarium

Prior to undergoing LASIK surgery with a WaveLight® Excimer Laser after surgery. One subject experienced retinal detachment, which was concluded to be unrelated to the treatment. A subject suffered from decreased vision in the treated eye, following blunt force trauma 4 days after surgery. Four of the eyes experienced transient or temporary decreases in vision prior to LASIK had the axis of astigmatism programmed as 115 degrees instead of the actual 155 degree angle.

The following complications were reported 6 months after LASIK: 0.8% (2/262) of the eyes had a retinal detachment or retinal vascular accident reported at the 3 month examination. The following complications were reported 6 months after LASIK: 0.2% (2/876) of the eyes had a lost, misplaced, or misaligned flap reported at the 1 month examination. The following complications were reported 6 months after LASIK: 0.9% (7/782) had an infection or significant dry eye that was unresponsive to treatment; 0.9% (7/782) had keratitis; 0.9% (7/782) had an unfavorable preoperative wavefront examination that precludes wave-front-guided treatment; and 3.4% (26/782) had a history of glaucoma.

The wavefront-guided LASIK procedure requires accurate and reliable data from the wavefront examination. Every step of every wavefront measurement that may be used as the basis for a wavefront-guided LASIK procedure must be validated by the user. Inaccurate or unreliable data from the wavefront examination will lead to an inaccurate treatment. Topography-guided LASIK requires preoperative topography maps of sufficient quality to use for planning a topography-guided LASIK treatment. Poor quality topography maps may affect the accuracy of the topography-guided LASIK treatment and may result in poor vision after topography-guided LASIK.

Precautions: The safety and effectiveness of the WaveLight® Excimer Laser Systems have not been established for patients with:

- myopia of any type
- hyperopia of any type
- astigmatism of any type
- previous corneal or intraocular surgery, or trauma in the ablation zone
- corneal abnormalities including, but not limited to, scars, irregular astigmatism and corneal warpage
- residual corneal thickness after ablation of less than 250 microns due to the increased risk for corneal ectasia
- pupil size below 7.0 mm after mydriatics which applied for wavefront-guided ablation planning
- history of glaucoma or ocular hypertension of > 23 mmHg
- taking the medications sumatriptan succinate (Imitrex®)
- corneal, lens and/or vitreous opacities including, but not limited to, cataract
- iritis, problems including, but not limited to, cold Kofoed and previous iris surgery compromising proper eye tracking
- taking medications likely to affect wound healing including (but not limited to) antidepressants.

In addition, safety and effectiveness of the WaveLight® Excimer Laser System have not been established for:

- treatments with an optical zone < 6.0 mm or > 6.5 mm in diameter, or an ablation zone > 9.0 mm in diameter
- wavefront-guided treatment targets different from emmetropia (plano) in which the wavefront calculated defocus (spherical term) has been adjusted
- In the WaveLight® Excimer Laser System clinical studies, there were few subjects with cylinder amounts > 4 D and ≤ 6 D. Not all complications, adverse events, and levels of effectiveness may have been determined for this population.

Pupil sizes should be evaluated under mesopic illumination conditions. Effects of treatment on vision under poor illumination cannot be predicted prior to surgery.

Adverse Events and Complications

Myopia: In the myopia clinical study, 0.2% (2/876) of the eyes had a lost, misplaced, or misaligned flap reported at the 1 month examination. The following complications were reported 6 months after LASIK: 0.9% (7/782) had an infection or significant dry eye that was unresponsive to treatment; 0.9% (7/782) had keratitis; 0.9% (7/782) had an unfavorable preoperative wavefront examination that precludes wave-front-guided treatment; and 3.4% (26/782) had a history of glaucoma.

Wavefront-Guided Myopia: The wavefront-guided myopia clinical study included 374 treated, 188 with wavefront-guided LASIK (Study Cohort) and 186 with Wavefront Optimize® LASIK (Control Cohort). In the Study Cohort and 166 of the Control Cohort were eligible to be followed at 6 months. In the Study Cohort, accountability at 1 month was 96.8%, at 3 months was 96.8%, and at 6 months was 93.3%. In the Control Cohort, accountability at 1 month was 94.6%, at 3 months was 94.6%, and at 6 months was 92.2%. The Ophthalmic Study Cohort that were eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better. Of the 166 eyes in the Study Cohort eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better. Of the 166 eyes in the Study Cohort eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better. Of the 166 eyes in the Study Cohort eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better.

In the Study Cohort, subjects who responded to a patient satisfaction questionnaire before and after LASIK reported the following visual symptoms at a "moderate" or "severe" level at least 1% higher at 3 months post-treatment than at baseline: sensitivity to light (53.9% vs. 43.3% at baseline); visual fluctuations (43.0% vs. 32.1% at baseline); and halos (42.3% vs. 37.0% at baseline).

Long term risk of LASIK for myopia with and without astigmatism have not been studied beyond 6 months.

Clinical Data

Myopia: The myopia clinical study included 901 eyes treated, of which 813 of 866 eligible eyes were followed for 12 months. Accountability at 3 months was 93.8%, at 6 months was 91.9%, and at 12 months was 93.9%. Of the 782 eyes that were eligible for the uncorrected visual acuity (UCVA) analysis of effectiveness at the 6-month stability time point, 98.3% were corrected to 20/40 or better, and 87.7% were corrected to 20/20 or better. Subjects who responded to a patient satisfaction questionnaire before and after LASIK reported the following visual symptoms at a "moderate" or "severe" level at least 1% higher at 3 months post-treatment than at baseline: visual fluctuations (28.6% vs. 12.8% at baseline).

Long term risks of LASIK for myopia with and without astigmatism have not been studied beyond 12 months.

Hyperopia: The hyperopia clinical study included 290 treated, of which 100 of 290 eligible eyes were followed for 12 months. Accountability at 3 months was 95.2%, at 6 months was 93.9%, and at 12 months was 69.9%. Of the 212 eyes that were eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 93.3% were corrected to 20/40 or better, and 69.4% were corrected to 20/20 or better. Subjects who responded to a patient satisfaction questionnaire before and after LASIK reported the following visual symptoms at a "moderate" or "severe" level at least 1% higher at 3 months post-treatment: halos (6.4%); visual fluctuations (6.1%); light sensitivity (4.9%); night driving glare (4.2%); and glare from bright lights (3.0%).

Long term risks of LASIK for hyperopia with and without astigmatism have not been studied beyond 12 months.

Topography-Guided Myopia: The topography-guided myopia clinical study included 374 treated, 188 with wavefront-guided LASIK (Study Cohort) and 186 with Wavefront Optimize® LASIK (Control Cohort). 166 of the Study Cohort and 166 of the Control Cohort were eligible to be followed at 6 months. In the Study Cohort, accountability at 1 month was 96.8%, at 3 months was 96.8%, and at 6 months was 93.3%. In the Control Cohort, accountability at 1 month was 94.6%, at 3 months was 94.6%, and at 6 months was 92.2%. Of the 166 eyes in the Study Cohort that were eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better. Of the 166 eyes in the Study Cohort eligible for the UCVA analysis of effectiveness at the 6-month stability time point, 99.4% were corrected to 20/40 or better, and 93.4% were corrected to 20/20 or better.

In the Study Cohort, subjects who responded to a patient satisfaction questionnaire before and after LASIK reported the following visual symptoms at a "moderate" or "severe" level at least 1% higher at 3 months post-treatment than at baseline: halos (45.4% vs. 36.6% at baseline); and visual fluctuations (21.9% vs. 18.3% at baseline).

Long term risks of wavefront-guided LASIK for myopia with and without astigmatism have not been studied beyond 6 months.

Topography-Guided Myopia: The topography-guided myopia clinical study included 249 treated, of which 230 eyes were followed for 12 months. Accountability at 3 months was 99.2%, at 6 months was 98.0%, and at 12 months was 92.4%. Of the 247 eyes that were eligible for the UCVA analysis at the 3-month stability time point, 99.2% were corrected to 20/40 or better, and 92.7% were corrected to 20/20 or better. Subjects who responded to a patient satisfaction questionnaire before and after LASIK reported the following visual symptoms as "marked" or "severe" at an incidence greater than 5% at 1 month after surgery: dryness (7% vs. 4% at baseline) and light sensitivity (7% vs. 5% at baseline). Visual symptoms continued to improve with time, and none of the visual symptoms were rated as being "marked" or "severe" with an incidence of at least 5% at 3 months or later after surgery.

Long term risks of topography-guided LASIK for myopia with and without astigmatism have not been studied beyond 12 months.

Information for Patients: Prior to undergoing LASIK surgery with a WaveLight® Excimer Laser System, prospective patients must receive a copy of the relevant Patient Information Booklet, and must be informed of the alternatives for correcting their vision, including (but not limited to) eyeglasses, contact lenses, photorefractive keratectomy, and other refractive surgeries.

Attention: Please refer to a current WaveLight® Excimer Laser System Procedure Manual for a complete listing of the indications, complications, warnings, precautions, and side effects.

* Trademarks are property of their respective owners.