The physical / optical properties of the lens are:

- **Specific Gravity:** 1.064
- **Refractive Index:** 1.426
- **Light Transmittance:** C.I.E. value - at least 95%
- **Water Content:** 36%
- **Oxygen Permeability:**
  - (Boundary Corrected) $91 \times 10^{-11} \text{cm}^2 \text{O}_2/(\text{sec} \times \text{cm}^2 \times \text{mmHg}) @ 35^\circ \text{C}$ Polarographic Method
  - (Non-Edge Corrected) $101 \times 10^{-11} \text{cm}^2 \text{O}_2/(\text{sec} \times \text{cm}^2 \times \text{mmHg}) @ 35^\circ \text{C}$ Polarographic Method

The PureVision Toric Contact Lenses, with AerGel™ technology lens material, are manufactured by a cast molding process and are surface treated by the Performa™ surface treatment process which transforms hydrophobic silicone to hydrophilic silicate. The anterior surface of the lens contains the aspheric optic zone, prism ballast and comfort chamfer feature of the PureVision Toric Contact Lens. The posterior surface is manufactured with a spherocylindrical curve to accommodate the required astigmatic power.

**CAUTION:** Federal (U.S.A.) law restricts this device to sale by or on the order of a licensed practitioner.

**IMPORTANT:**
This package insert and fitting guide has been developed to provide practitioners with information covering characteristics of the BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens with the Lo-Torque™ design and to illustrate fitting procedures. It is effective as of April 2005 and supersedes all prior fitting guides for the product described. Please read carefully and keep this information for future use.

This package insert and fitting guide is intended for the eye care professional, but should be made available to patients upon request. The eye care professional should provide the patient with the patient instructions that pertain to the patient’s prescribed lens and the recommended wearing schedule.

**DESCRIPTION:**
The BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens is a soft hydrophilic contact lens which is available as a flexible shell with a toric surface. The lens material, balafilcon A, is a copolymer of a silicone vinyl carbamate, N-vinyl-pyrrolidone, a siloxane crosslinker and a vinyl alanine wetting monomer, and is 36% water by weight when immersed in a sterile borate buffered saline solution. This lens is tinted blue with up to 300 ppm of Reactive Blue Dye 246.

**LENS PARAMETERS AVAILABLE**: The BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens is a hemispherical shell of the following dimensions:

- **Diameter:** 14.0mm
- **Center Thickness:** 0.05mm to 0.50mm
- **Base Curve:** 8.7mm
- **Sphere Powers:** -0.25D to -6.00D in 0.25D steps
- **Cylinder Powers:** -0.75D, -1.25D and -1.75D
- **Axis:** 10° to 180° in 10° Increments

*Additional powers may be introduced over time, check for product availability.

**Lens Prism:** Prism is located at the base of the lens to stabilize lens positioning when lens is on the eye.

**Comfort Chamfer:** A wedge-shaped tapered section on the anterior surface of the lens in the periphery of the lens from the 3 to 9 o’clock areas. This reduces lens thickness.

**HOW THE LENS WORKS (ACTIONS):**
In its hydrated state, the BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens has a unique Lo-Torque™ design that results in excellent stability and when placed on the cornea, acts as a refracting medium to focus light rays on the retina.

**INDICATIONS:**
The BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens is indicated for daily wear or extended wear from 1 to 30 days between removals, for cleaning and disinfection or disposal of the lens, as recommended by the eye care professional. The lens is indicated for the correction of refractive ametropia (myopia, hyperopia and astigmatism) in aphakic and/or not-aphakic persons with non-diseased eyes, exhibiting astigmatism of up to 5.00 diopters, that does not interfere with visual acuity. The lens may be prescribed for Frequent/Planned Replacement Wear or Disposable Wear in spherical powers ranging from +6.00D to -9.00D when prescribed for up to 30 days of extended wear and from +20.00D to -20.00D for daily wear or extended wear up to 7 days.
Note: See the WARNINGS reference to the relationship between lens wearing schedule and corneal complications.

FREQUENT/PLANNED REPLACEMENT WEAR
When prescribed for Frequent/Planned Replacement Wear, the PureVision Toric Contact Lens is to be cleaned, rinsed, and disinfected each time it is removed from the patient's eye and discarded after the recommended wearing period prescribed by the eye care professional. The lens may be disinfected using a chemical disinfection system.

DISPOSABLE WEAR
When prescribed for Disposable Wear, the PureVision Toric Contact Lens is to be discarded after each removal.

CONTRAINDICATIONS (REASONS NOT TO USE):
DO NOT USE the BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lens when any of the following conditions exist:
• Acute and subacute inflammation or infection of the anterior chamber of the eye
• Any eye disease, injury, or abnormality that affects the cornea, conjunctiva, or eyelids
• Severe insufficiency of lacrimal secretion (dry eyes)
• Corneal hypoesthesia (reduced corneal sensitivity)
• Any systemic disease that may affect the eye or be exaggerated by wearing contact lenses
• Allergic reactions of ocular surfaces or adnexa (surrounding tissue) that may be induced or exaggerated by wearing contact lenses or use of contact lens solutions
• Allergy to any ingredient, such as mercury or Thimerosal, in a solution which is to be used to care for the PureVision Toric Contact Lens
• Any active corneal infection (bacterial, fungal, or viral)
• If eyes become red or irritated

WARNINGS:
After a thorough eye examination, including appropriate medical background, patients should be fully apprised by the prescribing professional of all the risks with contact lens wear. Patients should be advised of the following warnings pertaining to contact lens wear:
• Problems with contact lenses and lens care products could result in serious injury to the eye. It is essential that patients follow their eye care professional's direction and all labeling instructions for proper use of lenses and lens care products, including the lens case. Eye problems, including corneal ulcers, can develop rapidly and lead to loss of vision.
• When prescribed for Frequent/Planned Replacement Wear, the need for strict compliance with the care regimen including cleaning of the lens case, wearing restrictions, wearing schedule, and follow-up visit schedule should be emphasized to the patient.
• Studies have shown that contact lens wearers who are smokers have a higher incidence of adverse reactions than nonsmokers.
• If a patient experiences eye discomfort, excessive tearing, vision changes, or redness of the eye, the patient should be instructed to immediately remove lenses and promptly contact his or her eye care professional.

EXTENDED WEAR
• The risk of microbial keratitis has been shown to be greater among users of extended wear contact lenses than among users of daily wear contact lenses. The risk among extended wear lens users increases with the number of consecutive days that the lenses are worn between removals, beginning with the first overnight use.

Some researchers believe that these complications are caused by one or more of the following: a weakening of the cornea's resistance to infections, particularly during a closed-eye condition, as a result of hypoxia; an eye environment which is somewhat more conducive to the growth of bacteria and other microorganisms, particularly when a regular periodic lens removal and disinfecting or disposal schedule has not been adhered to by the patient; improper lens disinfection or cleaning by the patient; contamination of lens care products; poor personal hygiene by the patient; patient unsuitability to the particular lens or wearing schedule; accumulation of lens deposits; damage to the lens; improper fitting; length of wearing time; and the presence of ocular debris or environmental contaminants.

While the great majority of patients successfully wear contact lenses, extended wear of lenses also is reported to be associated with a higher incidence and degree of epithelial microcysts and infiltrates, and endothelial polymegathism, which require consideration of discontinuation or restriction of extended wear. The epithelial conditions are reversible upon discontinuation of extended wear.

The risk of microbial keratitis has not been determined for this lens. Post marketing studies are in progress.

The reversibility of endothelial effects of contact lens wear has not been conclusively established. As a result, professionals views of extended wearing times vary from not prescribing extended wear at all to prescribing flexible wearing times from occasional overnight wear to prescribing extended wearing periods from 1 to 30 days with specified intervals of no lens wear for certain patients, with follow-up visits, and with proper care regimen.

PRECAUTIONS:
Precautions for Eye Care Professionals:
• Due to the small number of patients enrolled in clinical investigation of lenses, all refractive powers, design configurations, or lens parameters available in the lens material are not evaluated in significant numbers. Consequently, when selecting an appropriate lens design and parameters, the eye care professional should consider all characteristics of the lens that can affect lens performance and ocular health, including oxygen permeability, wettability, central and peripheral thickness, and optic zone diameter.

The oxygen transmissibility is below the established threshold required to prevent overnight corneal edema for portions of the power range, including plus powers and some low minus power lenses. In the U.S. clinical study of the PureVision (spherical) lens, the rate of infiltrative keratitis was found to be higher with higher lens powers (see Clinical Study section of the package insert).
• The potential impact of these factors on the patient's ocular health should be carefully weighed against the patient's need for refractive correction; therefore, the continuing ocular health of the patient and lens performance on eye should be carefully monitored by the prescribing eye care professional.
• Eye care professionals should instruct the patient to REMOVE A LENS IMMEDIATELY if an eye becomes red or irritated.
• Fluorescein, a yellow dye, should not be used while the lenses are on the eyes. The lenses absorb this dye and become discolored. Whenever fluorescein is used in eyes, the eyes should be flushed with sterile saline solution that is recommended for in-eye use.
• The patient should be instructed to always discard disposable lenses and lenses worn on a frequent/planed replacement schedule after the recommended wearing schedule prescribed by the eye care professional.
• Some patients will not be able to tolerate continuous wear even if able to tolerate the same or another lens on a daily wear basis. Some patients who are able to tolerate continuous wear will not be able to wear their lenses continuously for 30 days. Patients should be carefully evaluated for continuous wear prior to prescription and dispensing, and eye care professionals should conduct early and frequent follow-up examination to determine ocular response to continuous wear.
• As with any contact lens, follow-up visits are necessary to assure the continuing health of the patient’s eyes. The patient should be instructed as to a recommended follow-up schedule.
• Aphakic patients should not be fitted with PureVision Contact Lenses until the determination is made that the eye has healed completely.

Eye care professionals should carefully instruct patients about the following lens care and safety precautions. It is strongly recommended that patients be provided with a copy of the PureVision Toric Patient Information Booklet available from BAUSCH & LOMB® and understand its contents prior to dispensing the lenses.

Handling Precautions:
• Always wash and rinse hands before handling lenses. Do not get cosmetics, lotions, soaps, creams, deodorants, or sprays in the eyes or on the lenses. It is best to put on lenses before putting on makeup. Water-base cosmetics are less likely to damage lenses than oil-base products.
• Be sure that before leaving the eye care professional’s office, the patient is able to remove lenses promptly or have someone else available to remove them.
• Be certain that the fingers or hands are free of foreign materials before touching lenses, as microscopic scratches of the lenses may occur, causing distorted vision and/or injury to the eye.
• Always handle lenses carefully and avoid dropping them.
• Do not touch the lenses with fingernails.
• Carefully follow the handling, insertion, removal, cleaning disinfecting, storing and wearing instructions in the Patient Information Booklet for the PureVision Toric Contact Lenses and those prescribed by the eye care professional.
• Never use tweezers or other tools to remove lenses from the lens container unless specifically indicated for that use. Pour the lens into the hand.

Solution Precautions:
Do not use the Allergan® Ultracare® Disinfecting System or any of its components (Ultracare® Disinfecting Solution, Ultracare® Neutralizing Tablets, Lens Plus Daily Cleaner, and Ultrasyme® Enzymatic Cleaner) to clean and disinfect the PureVision Toric Contact Lens because the lens dimension will be altered.
• Always use fresh unexpired lens care solutions.
• Always follow directions in the package inserts for the use of contact lens solutions.
• Sterile unpreserved solutions, when used, should be discarded after the time specified in the labeling directions.
• Always keep the lenses completely immersed in the recommended storage solution when lenses are not being worn (stored). Prolonged periods of drying will damage lenses. Follow the lens care directions for Care for a Dried Out (Dehydrated) Lens in the patient information booklet if lens surface does become dried out.
• Do not use saliva or anything other than the recommended solution for lubricating or wetting lenses.
• Tap water, distilled water or homemade saline should not be used as a substitute for any component in the lens care regimen since they have been associated with an Acanthamoeba keratitis infection.

• Never use conventional hard contact lens solutions that are not also recommended for use with prescribed lenses.
• Do not mix or alternate lens care systems or solutions unless indicated in the lens care system labeling.
• Do not heat the chemical disinfection solution or lenses.

Lens Wearing Precautions:
• Never wear lenses beyond the period recommended by the eye care professional.
• If the lens sticks (stops moving) on the eye, follow the recommended directions on Care for a Sticking Lens. The lens should move freely on the eye for the continued health of the eye. If nonmovement of the lens continues, the patient should be instructed to immediately consult his or her eye care professional.
• Avoid, if possible, all harmful or irritating vapors and fumes while wearing lenses.
• If aerosol products are used while wearing lenses, exercise caution and keep eyes closed until the spray has settled.

Lens Case Precautions:
• Contact lens cases can be a source of bacterial growth. To prevent contamination and to help avoid serious eye injury, always empty and rinse the lens case with fresh, sterile rinsing solution and allow to air dry.
• Lens cases should be replaced at regular intervals as recommended by the lens case manufacturer or eye care professional.

Topics to Discuss with the Patient:
• As with any contact lens, follow-up visits are necessary to assure the continuing health of the eyes. The patient should be instructed as to a recommended follow-up schedule.
• Patients should be advised about wearing lenses during sporting and water related activities. Exposure to water while wearing contact lenses in activities such as swimming, water skiing and hot tubs may increase the risk of ocular infection including but not limited to Acanthamoeba keratitis.
• Always contact the eye care professional before using any medicine in the eyes.

Who Should Know That the Patient is Wearing Contact Lenses:
• Patients should inform their doctor (health care practitioner) about being a contact lens wearer.
• Patients should always inform their employer of being a contact lens wearer. Some jobs may require the use of eye protection equipment or may require that you do not wear lenses.

ADVERSE REACTIONS:
The patient should be informed that the following problems may occur:
• Eyes stinging, burning, itching (irritation), or other eye pain
• Comfort is less than when lens was first placed on eye
• Abnormal feeling of something in the eye (foreign body, scratched area)
• Excessive watering (tearing) of the eyes
• Unusual eye secretions
• Redness of the eyes
• Reduced sharpness of vision (poor visual acuity)
• Blurred vision, rainbows, or halos around objects
• Sensitivity to light (photophobia)
• Dry eyes

If the patient notices any of the above, he or she should be instructed to:
• Immediately remove the lenses.
If the discomfort or problem stops, then look closely at the lens. If the lens is in any way damaged, do not put the lens back on the eye. Place the lens in the storage case and contact the eye care professional. If the lens has dirt, an eyelash, or other foreign body on it, or the problem stops and the lens appears undamaged, the patient should thoroughly clean, rinse, and disinfect the lenses; then reinsert them. After reinsertion, if the problem continues, the patient should immediately remove the lenses and consult the eye care professional.

If the above symptoms continue after removal of the lens, or upon reinsertion of a lens, or upon insertion of a new lens, the patient should immediately remove the lenses and contact his or her eye care professional or physician, who must determine the need for examination, treatment or referral without delay. (See Important Treatment Information for Adverse Reactions.) A serious condition such as infection, corneal ulcer, corneal vascularization, or iritis may be present, and may progress rapidly. Less serious reactions such as abrasions, epithelial staining or bacterial conjunctivitis must be managed and treated carefully to avoid more serious complications.

Important Treatment Information for Adverse Reactions
Sight-threatening ocular complications associated with contact lens wear can develop rapidly, and therefore early recognition and treatment of problems are critical. Infectious corneal ulceration is one of the most serious potential complications, and may be ambiguous in its early stage. Signs and symptoms of infectious corneal ulceration include discomfort, pain, inflammation, purulent discharge, sensitivity to light, cells and flare and corneal infiltrates.

Initial symptoms of a minor abrasion and an early infected ulcer are sometimes similar. Accordingly, such epithelial defect, if not treated properly, may develop into an infected ulcer. In order to prevent serious progression of these conditions, a patient presenting symptoms of abrasions or early ulcers should be evaluated as a potential medical emergency, treated accordingly, and be referred to a corneal specialist when appropriate. Standard therapy for corneal abrasions such as eye patching or the use of steroids or steroid/antibiotic combinations may exacerbate the condition. If the patient is wearing a contact lens on the affected eye when examined, the lens should be removed immediately and the lens and lens care products retained for analysis and culturing.

CLINICAL STUDY:
The following clinical results are provided for informational purposes. It is important to note that the results below are from a study conducted with the PureVision Contact Lens which has the same lens material, but different lens design (spherical). The study was conducted with subjects not requiring astigmatic correction.

STUDY DESCRIPTION
Study Design
The objective of this 12-month study was to evaluate the safety and efficacy of the BAUSCH & LOMB® PureVision™ (balaficon A) Visibility Tinted Contact Lenses worn on a 30-day continuous wear basis, compared to a conventional Contact lens worn on a 7-day continuous wear basis. A total of 1640 eyes (820 subjects) were enrolled into this study. Subjects were fitted with a PureVision Contact Lens on one eye while the contralateral eye was fitted with a Control lens. Subjects were instructed to replace the PureVision Contact Lens with a new lens every 30 days, and to wear the Control lens overnight for up to six consecutive nights per week. Eyes had one night without lens wear after the scheduled removal. The Control lens was to be replaced with a new lens every 14 days.

Six hundred ten (610) subjects completed the one-year study. Ten subjects discontinued in the daily wear adaptation period, 182 subjects discontinued during the extended wear phase and 18 subjects were not dispensied lenses.

Patient Assessments
Subjects were evaluated at follow-up visits scheduled after 24 hours, 10 days, 1 month, 3 months, 6 months, 9 months, and 12 months of lens wear.

Demographics
Subject recruitment was open to adapted and unadapted contact lens wearers. There were no restrictions as to the subject’s gender or occupation, but subjects were required to be of legal age (typically 18 or 21) and have the legal capacity to volunteer. The ages of the subjects ranged from 18 to 74 years of age, with a mean age of 33.6, and included 574 females and 228 males, with a ratio of 2.52 females to every male. For the PureVision Contact Lens the power range used was –0.50D to –9.00D. For the Control lens the power range was –0.50D to –8.50D.

The previous lens wearing experience of the subjects that participated in the study was 5% no lens wear, 43% daily wear, and 51% continuous wear. The refractive errors of the subjects that participated in the study were from –0.25D to –11.75D, and included up to –2.00D of astigmatism.

SUMMARY OF DATA ANALYSES
Summary of Data Analyses
The key endpoints for this study were:
1. grade 2 and higher slit lamp findings (safety endpoint),
2. grade 2 and higher corneal infiltrates (safety endpoint), and
3. contact lens corrected visual acuity worse than 20/40 (efficacy endpoint).

For each key endpoint, the rates (incidents of endpoint/number of eyes) experienced by eyes in the PureVision Contact Lens and Control lenses were calculated. The difference in rates between the two lens types was determined and a 95% confidence interval for the difference was calculated. For each key endpoint a “clinically significant difference” in the rates was established before the study started. These “clinically significant differences” were as follows: 10% for total slit lamp findings ≥ Grade 2, 5% for corneal infiltrates ≥ Grade 2, and 5% for the acuity endpoint. For example, if the true rates of endpoint infiltrates in the subject population were 9.99% in the PureVision Contact Lens and 5% in the Control lens, these rates would be considered substantially equivalent (difference <5%).

In order to be successful for a given endpoint, the upper 95% confidence limit for the difference in the study rates had to be less than the pre-established “clinically significant difference.” This means that we are 95% confident that the true difference is within tolerance. The safety and efficacy goals were met for all three key endpoints. Results are as follows:

<table>
<thead>
<tr>
<th>Endpoint</th>
<th>PureVision</th>
<th>Control</th>
<th>Relative Risk PureVision/Control</th>
<th>Difference in %</th>
<th>Upper 95% Confidence Level</th>
<th>Clinically Significant Difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slit Lamp Findings ≥ Grade 2</td>
<td>138</td>
<td>139</td>
<td>17.5%</td>
<td>17.6%</td>
<td>1.0</td>
<td>-0.1%</td>
</tr>
<tr>
<td>Corneal Infiltrates ≥ Grade 2</td>
<td>23</td>
<td>10</td>
<td>2.9%</td>
<td>3.3%</td>
<td>2.3</td>
<td>1.6%</td>
</tr>
<tr>
<td>Visual Acuity Worse than 20/40</td>
<td>0</td>
<td>2</td>
<td>0.0%</td>
<td>0.3%</td>
<td>0.0</td>
<td>-0.3%</td>
</tr>
</tbody>
</table>
Summary of Slit Lamp Findings
Slit lamp examinations were conducted at every study visit. Each graded slit lamp parameter was scored on a qualitative grade scale ranging from 0 to 4, with Grade 0 representing the absence of findings, and Grades 1 through 4 representing successively worse findings. For each study eye, a determination was made for each parameter as to whether, or not a positive finding was presented at any visit. The following table describes slit lamp findings ≥ Grade 2 and ungraded slit lamp findings.

<table>
<thead>
<tr>
<th>Graded Slit Lamp Findings (≥ Grade 2)</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any Finding1,2</td>
<td>17.5%</td>
<td>17.6%</td>
</tr>
<tr>
<td>Corneal Staining</td>
<td>8.2%</td>
<td>8.4%</td>
</tr>
<tr>
<td>Limbal Injection</td>
<td>3.7%</td>
<td>4.3%</td>
</tr>
<tr>
<td>Bulbar Injection</td>
<td>5.2%</td>
<td>4.7%</td>
</tr>
<tr>
<td>Tarsal Conjunctival Abnormalities</td>
<td>3.9%</td>
<td>3.9%</td>
</tr>
<tr>
<td>Corneal Infiltrates1</td>
<td>2.9%</td>
<td>1.3%</td>
</tr>
<tr>
<td>Epithelial Edema</td>
<td>1.3%</td>
<td>1.4%</td>
</tr>
<tr>
<td>Epithelial Microcysts</td>
<td>1.0%</td>
<td>1.0%</td>
</tr>
<tr>
<td>Corneal Neovascularization</td>
<td>1.0%</td>
<td>1.7%</td>
</tr>
</tbody>
</table>

Ungraded Slit Lamp Findings

<table>
<thead>
<tr>
<th>Other Anterior Segment Abnormalities3</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>13.2%</td>
<td>13.8%</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>External Adnexa Abnormalities</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7%</td>
<td>2.7%</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Conjunctivitis</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.4%</td>
<td>2.0%</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Corneal Striae</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0%</td>
<td>0.3%</td>
<td></td>
</tr>
</tbody>
</table>

1/ Slit Lamp Finding and Corneal Infiltrates ≥ Grade 2 were the safety endpoints for this study.
2/ The total of all Graded slit lamp findings does not equal the category of Any Finding.
3/ The more common findings identified as Other Anterior Segment Abnormalities included: conjunctival staining; dimple veils; mucin balls; lipid deposits; and ghost vessels.

It should be noted that the PureVision Contact Lens and Control lenses were each fit on only the right or left eye for each subject. Rates per subject are expected to be higher when lenses are fit on both eyes.

Corneal Infiltrates

The following table describes the rate of corneal infiltrates according to the lens power used.

<table>
<thead>
<tr>
<th>Lens Power</th>
<th>Corneal Infiltrates (≥ Grade 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PureVision</td>
<td></td>
</tr>
<tr>
<td>Plano to -3.00</td>
<td>1.7%</td>
</tr>
<tr>
<td>-3.25 to -6.00</td>
<td>3.2%</td>
</tr>
<tr>
<td>&gt;-6.00</td>
<td>6.4%</td>
</tr>
<tr>
<td>Total</td>
<td>2.9%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Control</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Plano to -3.00</td>
<td>0.9%</td>
</tr>
<tr>
<td>-3.25 to -6.00</td>
<td>1.5%</td>
</tr>
<tr>
<td>&gt;-6.00</td>
<td>1.3%</td>
</tr>
<tr>
<td>Total</td>
<td>1.3%</td>
</tr>
</tbody>
</table>

Other Lens-Related Adverse Events

In addition to the outcomes described above, the following lens related adverse events were noted. This table does not include conjunctivitis or tarsal conjunctival abnormalities, e.g., giant papillary conjunctivitis.

<table>
<thead>
<tr>
<th>Other Important Lens-Related Adverse Events</th>
<th>PureVision</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corneal Scar</td>
<td>14 (1.8%)</td>
<td>5 (0.6%)</td>
</tr>
<tr>
<td>Other Ocular Inflammation*</td>
<td>10 (1.3%)</td>
<td>2 (0.3%)</td>
</tr>
<tr>
<td>Anterior Chamber Reaction</td>
<td>2 (0.3%)</td>
<td>1 (0.1%)</td>
</tr>
<tr>
<td>Permanant Loss of Vision</td>
<td>0 (0.0%)</td>
<td>0 (0.0%)</td>
</tr>
</tbody>
</table>

* Other Ocular Inflammation includes episcleritis, scleritis, iritis/uveitis.

Efficacy Outcomes

The contact lens visual acuity was measured at each scheduled and unscheduled follow-up visit throughout the one-year study. For the 610 subjects that completed the study, visual acuity of 20/20 or better was reported for 87% and 86% of the measurements for the PureVision Contact Lens and Control lens, respectively. Similarly, visual acuity of 20/25 or better was reported 98% and 97% of the times for the PureVision Contact Lens and Control lens.

Wearing Time

In this U.S. clinical study subjects were required to maintain a minimum wearing time in order to continue in the study. For the subjects that completed the study, the average continuous wear time for the PureVision Contact Lens was at least 28.0 days per month, from the 2-Month visit through the 12-Month visit. At these visits the same subjects reported they were able to wear the PureVision Contact Lens at least 22 days continuously with 94% of the times they were asked.

During the course of the study, 15 subjects were discontinued from the study because they were not able to wear the PureVision Contact lens for 30 days. Twenty-one (21) subjects were discontinued from the study because they were not able to wear the Control lens for 7 days.

Overnight Corneal Swelling

A study was conducted to assess the corneal swelling response induced by overnight contact lens wear. Twenty-four (24) subjects each wore either a -3.00 – 0.75 x 180° PureVision Toric Contact Lens (Test Lens) or a -3.00D PureVision Contact Lens (Control lens) on the contralateral eye overnight under closed eye conditions for approximately eight hours. The corneal swelling, measured as the percent increase in the center thickness of the cornea, of the eyes wearing a PureVision Toric Lens (4.1%) was compared to the swelling response to the Control lens (3.6 %). The responses were not statistically different (p-value > 0.20).

SELFICATION OF PATIENTS:
The eye care professional should not fit patients who cannot or will not adhere to a recommended care or replacement regimen, or are unable to place and remove the lenses should not be provided with them.

Failure to follow handling and cleaning instructions could lead to serious eye infections which might result in corneal ulcers.

Patient communication is vital because it relates not only to patient selection but also to ensure compliance. It is also necessary to discuss the information contained in the Patient Information Booklet with the patient at the time of the initial examination.
Patients selected to wear PureVision Toric Contact Lenses should be chosen for their motivation to wear contact lenses, general health and cooperation. The eye care professional must take care in selecting, examining and instructing contact lens patients. Patient hygiene and willingness to follow practitioner instructions are essential to their success.

A detailed history is crucial to determining patient needs and expectations. Your patient should be questioned regarding vocation, desired lens wearing time (full or part time), and desired lens usage (reading, recreation or hobbies).

Initial evaluation of the trial lens should be preceded by a complete eye examination, including visual acuity with and without correction at both distance and near, keratometry and slit lamp examination.

It is normal for the patient to experience mild symptoms such as lens awareness, variable vision, occasional tearing (watery eyes) and slight redness during the adaptation period. Although the adaptation period varies for each individual, generally within one week these symptoms will disappear. If these symptoms persist, the patient should be instructed to contact his or her eye care professional.

FITTING PROCEDURE:

1. Pre-Fitting Examination

   A pre-fitting patient history and examination are necessary to:
   • determine whether a patient is a suitable candidate for daily wear contact lenses (consider patient hygiene and mental and physical state),
   • make ocular measurements for initial contact lens parameter selection, and
   • collect and record baseline clinical information to which post-fitting examination results can be compared.

   A prefitting examination should include sphero-cylinder refraction and VA, keratometry, and biomicroscopic examination.

2. Initial Lens Power Selection Selection

   a. Select the initial trial lens from the Toric Diagnostic Lens Set with a power most similar to the patients refractive needs, or order a diagnostic lens to the prescription which most closely matches that of the patient.

   b. Place the lens on the eye and allow the lens to remain on the eye long enough (10 to 20 minutes) to achieve a state of equilibrium. Small variations in the tonicity, pH of the lens solutions, and individual tear composition may cause slight changes in fitting characteristics.

   c. Allow any increase in tear flow to subside before evaluating the lens. The time required will vary with the individual.

3. Initial Lens Evaluation

   A. To determine proper lens parameters, observe the lens relationship to the eye using a slit lamp. The toric diagnostic lens is used to optimize lens fitting characteristics and determine axis orientation. Lens power is determined by the spectacle refraction.

       • Rotation evaluation: The center guide mark should locate at the inferior limbus. The additional guide marks at 30° on either side can be used as reference points, and to help the professional assess axis orientation and stability. Once oriented, rotational rocking should be limited to less than 5°.

       • Movement: The lens should provide discernible movement with:

           - Primary gaze blink
           - Upgaze blink
           - Upgaze lag

       • Centration: The lens should provide full corneal coverage.

   B. Determine contact lens power. When the toric diagnostic lens does not have a power equivalent to their spectacle Rx, sphero-cylinder over-refractions will often be inaccurate and confusing. Therefore it is usually preferable to use the spectacle Rx as the only basis for the contact lens power. The sphere and cylinder power of the spectacle Rx becomes the sphere and cylinder power of the contact lens. There are two exceptions:

      1. If spectacle cylinder power falls between available contact lens cylinder powers, prescribe the lesser contact lens cylinder power. The sphere power can be increased -0.25D to compensate if desired. Of course, this can vary depending on your interpretation of the patient's subjective responses.

      Example:

      Spectacle Rx -2.00 -1.00 X 180
      Contact Lens Power Ordered -2.25 -0.75 X 180

      2. When the spectacle lens power in any principle meridian is greater than 4.00D, the spectacle refraction should be vertexed to the corneal plane. This can affect both the sphere and cylinder powers ordered.

      Example:

      Spectacle Rx -5.00 -2.75 X 180
      Contact Lens Power Ordered -4.75 -2.25 X 180

   C. Determine contact lens axis. Note the orientation of the guide marks relative to the vertical meridian. Regardless of which eye the lens is on, if the rotation is clockwise but stable, note the amount of rotation, add it to the refractive cylinder axis and order the resulting axis. If the rotation has stabilized counter-clockwise, again note the rotation, subtract it from the refractive axis and order the resulting axis. The guide marks can be used to help you calculate the axis of the desired Rx lens.

   For Example:

      Spectacle Rx -2.50 -1.25 X 80
      Rotation 20° clockwise
      Final Lens Prescription: -2.50 -1.25 X 100

   D. Select patient's lenses.

   E. Evaluate orientation of final Rx lenses. The orientation of the prescription should be the same as that observed for the Fitting Set Lenses. For example, if the lens rotated clockwise 15° then the final prescription lens should also rotate clockwise 15°.

4. Criteria of a Well-Fitted Lens

   If the initial lens selection fully covers the cornea, provides discernible movement after a blink, is comfortable for the patient and provides satisfactory visual performance, it is a well fitted lens and can be dispensed.

5. Characteristics of a Tight (Steep) Lens

   A lens which is much too steep may subjectively and objectively cause distortion which will vary after a blink. However, if a lens is only marginally steep, the initial subjective and objective vision and comfort findings may be quite good. A marginally steep lens may be differentiated from a properly fitted lens by having the patient gaze upward. A properly fitted lens will tend to slide downward approximately 0.5mm while a steep lens will remain relatively stable in relationship to the cornea, particularly with the blink.

   With your finger, gently rotate the lens approximately 45° to the temporal side. It should reorient within 5 to 10 blinks back to the same stabilized position.
6. Characteristics of a Loose (Flat) Lens
If the lens is too flat, it will:
- Decenter, especially on post-blink.
- Have a tendency to edge lift inferiorly and sit on the lower lid, rather than positioning between the sclera and palpebral conjunctiva.
- Have a tendency to be uncomfortable and irritating with fluctuating vision.
- Have a tendency to drop or lag greater than 2.0mm on upgaze post-blink.

7. Follow-up Care
a. Follow-up examinations are necessary to ensure continued successful contact lens wear. From the day of dispensing, the following schedule is a suggested guideline for follow up.
   - 24 hours
   - 10 days
   - 1 month
   - 3 months
   - every six months thereafter

   At the initial follow-up evaluations the eye care professional should again reassure the patient that any of the previously described adaptive symptoms are normal, and that the adaptation period should be relatively brief. Depending on the patients prior experience with contact lenses and/or continuous wear, the eye care professional may consider prescribing a one week period of daily wear adaption prior to beginning continuous wear.

b. Prior to a follow-up examination, the contact lenses should be worn for at least 4 continuous hours and the patient should be asked to identify any problems which might be occurring related to contact lens wear. If the patient is wearing the lenses for continuous wear, the follow-up examination should be conducted as early as possible the morning after overnight wear.

c. With lenses in place on the eyes, evaluate fitting performance to assure that CRITERIA OF A WELL FITTED LENS continue to be satisfied. Examine the lenses closely for surface deposition and/or damage.

d. After the lens removal, instill sodium fluorescein [unless contraindicated] into the eyes and conduct a thorough biomicroscopy examination.
   1. The presence of vertical corneal striae in the posterior central cornea and/or corneal neovascularization may be indicative of excessive corneal edema.
   2. The presence of corneal staining and/or limbal-conjunctival hyperemia can be indicative of an unclean lens, a reaction to solution preservatives, excessive lens wear, and/or a poorly fitting lens.
   3. Papillary conjunctival changes may be indicative of an unclean and/or damaged lens.

   If any of the above observations are judged abnormal, various professional judgments are necessary to alleviate the problem and restore the eye to optimal conditions. If the CRITERIA OF A WELL FITTED LENS are not satisfied during any follow-up examination, the patient should be re-fitted with a more appropriate lens.

PRACTITIONER FITTING SETS:
Lenses must be discarded after each use and must not be used from patient to patient.

WEARING SCHEDULE:
The wearing and replacement schedules should be determined by the eye care professional. Regular checkups, as determined by the eye care professional, are extremely important.

Continuous Wear (Greater than 24 hours or while asleep):
The wearing schedule should be determined by the prescribing eye care professional for each individual patient, based upon a full examination and patient history as well as the practitioner’s experience and professional judgment. Bausch & Lomb recommends beginning continuous wear patients with the recommended initial daily wear schedule, followed by a period of daily wear, and then gradual introduction of continuous wear one night at a time, unless individual considerations indicate otherwise.

The practitioner should examine the patient in the early stages of continuous wear to determine the corneal response. The lens must be removed, cleaned and disinfected or disposed of and replaced with a new lens, as determined by the prescribing eye care professional. (See the factors discussed in the Warnings section.) Once removed, a lens should remain out of the eye for a period of rest overnight or longer, as determined by the prescribing eye care professional.

MONOVISION FITTING GUIDELINES:

1. Patient Selection

   A. Monovision Needs Assessment
   For a good prognosis the patient should have adequately corrected distance and near visual acuity in each eye. The amblyopic patient or the patient with significant astigmatism (greater than one [1] diopter) in one eye may not be a good candidate for monovision with the PureVision Toric Contact Lenses.

   Occupational and environmental visual demands should be considered. If the patient requires critical vision (visual acuity and stereopsis) it should be determined by trial whether this patient can function adequately with monovision. Monovision contact lens wear may not be optimal for such activities as:
   (1) visually demanding situations such as operating potentially dangerous machinery or performing other potentially hazardous activities; and
   (2) driving automobiles (e.g., driving at night). Patients who cannot pass their state drivers license requirements with monovision correction should be advised to not drive with this correction, OR may require that additional over-correction be prescribed.

   B. Patient Education
   All patients do not function equally well with monovision correction. Patients may not perform as well for certain tasks with this correction as they have with bifocal reading glasses. Each patient should understand that monovision can create a vision compromise that may reduce visual acuity and depth perception for distance and near tasks. During the fitting process it is necessary for the patient to realize the disadvantages as well as the advantages of clear near vision in straight ahead and upward gaze that monovision contact lenses provide.
2. Eye Selection
   A. Ocular Preference Determination Methods
      Generally, the non-dominant eye is corrected for near vision. The
      following test for eye dominance can be used.

      Method 1 - Determine which eye is the "sighting dominant eye." Have
      the patient point to an object at the far end of the room. Cover one
      eye. If the patient is still pointing directly at the object, the eye being
      used is the dominant (sighting) eye.

      Method 2 - Determine which eye will accept the added power with the
      least reduction in vision. Place a trial spectacle near add lens in front of
      one eye and then the other while the distance refractive error correction
      is in place for both eyes. Determine whether the patient functions best
      with the near add lens over the right or left eye.

   B. Refractive Error Method
      For anisometropic corrections, it is generally best to fit the more
      hyperopic (less myopic) eye for distance and the more myopic (less
      hyperopic) eye for near.

   C. Visual Demands Method
      Consider the patient's occupation during the eye selection process
      to determine the critical vision requirements. If a patient's gaze
      for near tasks is usually in one direction correct the eye on that
      side for near.

      Example:
      A secretary who places copy to the left side of the desk will usually
      function best with the near lens on the left eye.

3. Special Fitting Considerations
   Unilateral Lens Correction
   There are circumstances where only one contact lens is required. As
   an example, an emmetropic patient would only require a near lens
   while a bilateral myope may require only a distance lens.

   Example:
   A presbyopic emmetropic patient who requires a +1.75 diopter add
   would have a +1.75 lens on the near eye and the other eye left with-
   out a lens.

   A presbyopic patient requiring a +1.50 diopter add who is -2.50
   diopters myopic in the right eye and -1.50 diopters myopic in the left
   eye may have the right eye corrected for distance and the left uncor-
   rected for near.

4. Near Add Determination
   Always prescribe the lens power for the near eye that provides opti-
   mal near acuity at the midpoint of the patient's habitual reading dis-
   tance. However, when more than one power provides optimal reading
   performance, prescribe the least plus (most minus) of the powers.

5. Trial Lens Fitting
   A trial fitting is performed in the office to allow the patient to experi-
   ence monovision correction. Lenses are fit according to the directions
   in the general fitting guidelines.

   Case history and standard clinical evaluation procedure should be used
   to determine the prognosis. Determine which eye is to be corrected for
   distance and which eye is to be corrected for near. Next determine the
   near add. With trial lenses of the proper power in place observe the
   reaction to this mode of correction.

   Immediately after the correct power lenses are in place, walk across
   the room and have the patient look at you. Assess the patient's reaction
to distance vision under these circumstances. Then have the patient
look at familiar near objects such as a watch face or fingernails. Again
assess the reaction. As the patient continues to look around room at
both near and distance objects, observe the reactions. Only after these
vision tasks are completed should the patient be asked to read print.
Evaluate the patient's reaction to large print (e.g. typewritten copy) at
first and then graduate to news print and finally smaller type sizes.

   After the patient's performance under the above conditions are com-
   pleted, tests of visual acuity and reading ability under conditions of
   moderately dim illumination should be attempted.

   An initial unfavorable response in the office, while indicative of a
   guarded prognosis, should not immediately rule out a more extensive
   trial under the usual conditions in which a patient functions.

6. Adaptation
   Visually demanding situations should be avoided during the initial
   wearing period. A patient may at first experience some mild blurred
   vision, dizziness, headaches, and a feeling of slight imbalance. You
   should explain the adaptational symptoms to the patient. These
   symptoms may last for a brief minute or for several weeks. The
   longer these symptoms persist, the poorer the prognosis for successful
   adaptation.

   To help in the adaptation process the patient can be advised to first
   use the lenses in a comfortable familiar environment such as in the
   home.

   Some patients feel that automobile driving performance may not be
   optimal during the adaptation process. This is particularly true when
   driving at night. Before driving a motor vehicle, it may be recommend-
   ed that the patient be a passenger first to make sure that their vision is
   satisfactory for operating an automobile. During the first several weeks
   of wear (when adaptation is occurring), it may be advisable for the
   patient to only drive during optimal driving conditions. After adaptation
   and success with these activities, the patient should be able to drive
   under other conditions with caution.

7. Other Suggestions
   The success of the monovision technique may be further improved by
   having your patient follow the suggestions below.
   - Having a third contact lens (distance power) to use when critical
distance viewing is needed.
   - Having a third contact lens (near power) to use when critical near
viewing is needed.
   - Having supplemental spectacles to wear over the monovision
contact lenses for specific visual tasks may improve the success of
monovision correction. This is particularly applicable for those
patients who cannot meet state licensing requirements with a
monovision correction.
   - Make use of proper illumination when carrying out visual tasks.

   Success in fitting monovision can be improved by the following sugges-
tions.
   - Reverse the distance and near eyes if a patient is having trouble
adaptating.
   - Refine the lens powers if there is trouble with adaptation. Accurate
lens power is critical for presbyopic patients.
   - Emphasize the benefits of the clear near vision in straight ahead
and upward gaze with monovision.

   * The decision to fit a patient with a monovision correction is most
appropriately left to the eye care professional in conjunction with the
patient after carefully considering the patient's needs.
* All patients should be supplied with a copy of the PureVision Toric Contact Lens Patient Information Booklet.

HANDLING OF LENSES:
Patient Lens Care Directions:
When lenses are dispensed, the patient should be provided with appropriate and adequate instructions and warnings for lens care handling. The eye care professional should recommend appropriate and adequate procedures and products for each individual patient in accordance with the particular lens wearing schedule and care system selected by the practitioner, the specific instructions for such products and the particular characteristics of the patient.

Frequent/Planned Replacement: For complete information concerning the care, cleaning and disinfection of contact lenses refer to the PureVision Toric (balafilcon A) Visibility Tinted Contact Lens Patient Information Booklet.

Disposable Wear: For complete information concerning emergency lens care, refer to the PureVision Toric Contact Lens Patient Information Booklet.

CARE FOR A STICKING (NONMOVING) LENS:
If the lens sticks (stops moving), the patient should be instructed to use a lubricating or rewetting solution in their eye. The patient should be instructed to not use plain water, or anything other than the recommended solutions. The patient should be instructed to contact the eye care professional if the lens does not begin to move upon blinking after several applications of the solution, and to not attempt to remove the lens except on the advice of the eye care professional.

REPORTING OF ADVERSE REACTIONS:
All serious adverse experiences and adverse reactions observed in patients wearing BAUSCH & LOMB® PureVision™ Toric (balafilcon A) Visibility Tinted Contact Lenses or experienced with the lenses should be reported to:
Bausch & Lomb Incorporated
Rochester, New York 14609

Toll Free Telephone Number
In the Continental U.S., Alaska, Hawaii
1-800-828-9030
In New York State
1-800-462-1720
In Canada
1-888-459-5000

HOW SUPPLIED:
Each sterile lens is supplied in a plastic package containing borate buffered saline solution. The container is marked with the manufacturing lot number of the lens, the base curve, sphere power, cylinder power, axis, diameter and expiration date. Store lenses at room temperature (60°F - 80°F, 15°C - 25°C).

BAUSCH & LOMB INCORPORATED
Rochester, NY 14609

© Bausch & Lomb Incorporated.
All rights reserved worldwide.
Bausch & Lomb, Performa, AerGel and PureVision are trademarks of Bausch & Lomb Incorporated.
Other brand names/product names are trademarks of their respective owners.
U.S. Patents and other patents.
Printed in U.S.A.

Symbol Reference Guide for label and cartons:

CE 0050 Quality System Certification Symbol

Fee Paid for Waste Management

STERILE Sterile Using Steam or Dry Heat

See Instruction Leaflet

Storage Temperature

DIA ør Diameter

EXP( ) Use By Date (expiration date)

LOT Batch Code

PWR (Fv') Dioptr (lens power)

EC REP Authorized representative in European Community

Rx Only Caution: Federal law restricts this device to sale by or on the order of a licensed professional.

BC Base Curve

SPH Sphere power (diopters)

AX Cylinder Axis (degrees)

CYL Cylinder power (diopters)