

PROFESSIONAL FITTING GUIDE

FluoroPerm[®] 92 (paflucocon A)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear

FluoroPerm[®] 60 (paflucocon B)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear

FluoroPerm[®] 30 (paflucocon C)

Rigid Gas Permeable Contact Lenses for Daily Wear

FluoroPerm[®] 151 (paflucocon D)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear



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CAUTIONS: Federal (US) law restricts this device to sale by, or on the order of a licensed practitioner.
Nonsterile. Clean and condition lenses prior to use.

PRODUCT DESCRIPTIONS

FluoroPerm[®] 92 (paflucocon A) and FluoroPerm[®] 60 (paflucocon B) rigid gas permeable contact lenses for daily wear are available as lathe cut or molded contact lenses with spherical, aspheric, bifocal or toric anterior and/or posterior; or, bitoric surfaces in clear and tinted versions. FluoroPerm 92 and FluoroPerm 60 rigid gas permeable contact lenses for extended wear are available as lathe cut or molded contact lenses with spherical or aspheric anterior or posterior surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 92 and FluoroPerm 60 rigid gas permeable contact lens materials are both thermoset copolymers derived from fluorosilicone acrylate monomers.

The FluoroPerm 92 and FluoroPerm 60 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM[®] 92 (paflucocon A)

Refractive Index	1.453 (Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	93%
Luminous Transmittance (Green)	95%
Wetting Angle (Receding Angle) ⁺⁺	16°
Wetting Angle ⁺⁺⁺	64°
Specific Gravity	1.10
Hardness (Shore D)	81
Water Content	<1%
Oxygen Permeability [*]	92 x 10 ⁻¹¹ Dk at 35°C
Oxygen Permeability ^{**}	64 x 10 ⁻¹¹ Dk at 35°C

⁺Determination of the Spectral and Luminous Transmittance, ISO 8599: 1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺ Sessile Drop Technique per ANSI Z80.20, 8.11

^{*}(cm²/sec)(mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

^{**}(cm²/sec)(mL O₂) / (mL x mm Hg) ISO/ANSI Method, ISO 9913-1

FLUOROPERM[®] 60 (paflucocon B)

Refractive Index	1.453 (Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	95%
Luminous Transmittance (Crystal Blue)	98%
Luminous Transmittance (Green)	95%
Luminous Transmittance (Brown)	85%
Wetting Angle (Receding Angle) ⁺	14.7°
Wetting Angle ⁺⁺	62°
Specific Gravity	1.15
Hardness (Shore D)	83
Water Content	<1%
Oxygen Permeability [*]	60 x 10 ⁻¹¹ Dk at 35°C
Oxygen Permeability ^{**}	43 x 10 ⁻¹¹ Dk at 35°C

⁺Determination of the Spectral and Luminous Transmittance, ISO 8599: 1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

+++ Sessile Drop Technique per ANSI Z80.20, 8.11

*(cm²/sec)(mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec)(mL O₂) / (mL x mm Hg) ISO/ANSI Method, ISO 9913-1

FluoroPerm[®] 30 (paflucocon C) rigid gas permeable contact lenses for daily wear are available as lathe cut or molded contact lenses with spherical, aspheric, bifocal or toric anterior and/or posterior; or, bitoric surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 30 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

The FluoroPerm 30 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM[®] 30 (paflucocon C)

Refractive Index	1.466 (Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	94%
Luminous Transmittance (Crystal Blue)	98%
Luminous Transmittance (Majestic Blue)	79%
Luminous Transmittance (Green)	95%
Luminous Transmittance (Gray)	91%
Wetting Angle (Receding Angle) ⁺	12.8°
Wetting Angle ⁺⁺	61°
Specific Gravity	1.14
Hardness (Shore D)	84
Water Content	<1%
Oxygen Permeability [*]	30 x 10 ⁻¹¹ Dk at 35°C
Oxygen Permeability ^{**}	30 x 10 ⁻¹¹ Dk at 35°C

⁺Determination of the Spectral and Luminous Transmittance, ISO 8599: 1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

+++ Sessile Drop Technique per ANSI Z80.20, 8.11

*(cm²/sec)(mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec)(mL O₂) / (mL x mm Hg) ISO/ANSI Method, ISO 9913-1

FluoroPerm[®] 151 (paflucocon D) rigid gas permeable contact lenses for daily wear and extended wear are available as lathe cut or molded contact lenses with spherical front and back surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 151 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

The FluoroPerm 151 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM® 151 (paflucocon D)

Refractive Index	1.442 (Nd at 25°C)
Luminous Transmittance ⁺ (Blue)	93%
Luminous Transmittance (Crystal Blue)	98%
Wetting Angle (Receding Angle) ⁺	42°
Wetting Angle ⁺⁺	70°
Specific Gravity	1.10
Hardness (Shore D)	79
Water Content	<1%
Oxygen Permeability [*]	151 x 10 ⁻¹¹ Dk at 35°C
Oxygen Permeability ^{**}	100 x 10 ⁻¹¹ Dk at 35°C

⁺Determination of the Spectral and Luminous Transmittance, ISO 8599: 1994

⁺⁺Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺Sessile Drop Technique per ANSI Z80.20, 8.11

^{*}(cm²/sec)(mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

^{**}(cm²/sec)(mL O₂) / (mL x mm Hg) ISO/ANSI Method, ISO 9913-1

TINTS (not all colors available in all materials)

FluoroPerm rigid gas permeable contact lenses are available in nontinted (clear) and tinted (blue, crystal blue, majestic blue, gray, brown and green) versions. The tinted lenses contain one or more of the following color additives: D&C Green No. 6, Peroxide Yellow No. 9, D&C Violet No. 2 and D&C Red No. 17.

UV ABSORBER (not available in all colors and materials)

FluoroPerm rigid gas permeable contact lenses are available with an ultraviolet absorber. The ultraviolet absorber, Uvinul D-49, has been integrated as an additive within the FluoroPerm polymer matrix, blocking up to 97% of light below 380 nm. The UV absorber is 2,2'-dihydroxy-4,4'-dimethoxybenzophenone.

See Package Insert for light transmission comparison graph.

WARNING: UV-absorbing contact lenses are not substitutes for protective UV-absorbing eyewear such as UV-absorbing goggles or sunglasses. Persons should continue to use their protective UV-absorbing eyewear as directed.

NOTE: The effectiveness of wearing UV-absorbing contact lenses in preventing or reducing the incidence of ocular disorders associated with exposure to UV-light has not been established at this time.

LENS PARAMETERS (not all available in all materials)

Chord Diameter	7.0 to 11.5 mm
Center Thickness	0.05 to 0.70 mm
Base Curve	6.50 to 9.00 mm
Power Daily Wear	-20.00 to +12.00 Diopters
Power Extended Wear	-20.00 to + 8.00 Diopters
Bifocal Add Power	+0.25 to + 4.00 Diopters
Monocentric Bifocal Add Diameter	4.0 to 9.0 mm
Monocentric Bifocal Prism	1.0 to 2.5 Diopters
Concentric Bifocal Add Diameter	2.0 to 4.0 mm

ACTIONS

See Package Insert (Actions) for the actions of each product; FluoroPerm[®] 92 (paflucocon A), FluoroPerm[®] 60 (paflucocon B), FluoroPerm[®] 30 (paflucocon C) and FluoroPerm[®] 151 (paflucocon D).

INDICATIONS

Device Name: FluoroPerm[®] 30 (paflucocon C) rigid gas permeable contact lenses

FluoroPerm[®] 30 rigid gas permeable contact lenses are indicated for daily wear as recommended by the eye care practitioner.

FluoroPerm[®] 30 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 30 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes. FluoroPerm[®] 30 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 30 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

Device Name: FluoroPerm[®] 60 (paflucocon B) rigid gas permeable contact lenses

FluoroPerm[®] 60 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm[®] 60 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm[®] 60 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 60 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm[®] 60 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 60 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

Device Name: FluoroPerm[®] 92 (paflucocon A) rigid gas permeable contact lenses

FluoroPerm[®] 92 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm[®] 92 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm[®] 92 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 92 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm[®] 92 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 92 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

Device Name: FluoroPerm[®] 151 (paflucocon D) rigid gas permeable contact lenses

FluoroPerm[®] 151 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm[®] 151 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm[®] 151 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 151 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm[®] 151 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 151 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

CONTRAINDICATIONS, WARNINGS, PRECAUTIONS AND ADVERSE REACTIONS

See Package Insert [Contraindications (Reasons Not To Use)] and (Warning) for each product; FluoroPerm[®] 92 (paflucocon A), FluoroPerm[®] 60 (paflucocon B) and FluoroPerm[®] 151 (paflucocon D).

See Package Insert [Contraindications (Reasons Not To Use)] and (Wearing Schedule) for FluoroPerm[®] 30 (paflucocon C).

See Package Insert, General Information (Warnings), (Precautions) and [Adverse Effects (Problems And What To Do)] for all products.

LENS HANDLING

CAUTION: Nonsterile. Clean and condition lenses prior to use.

1. Prior to fitting, wash your hands and rinse them thoroughly to remove all traces of soap.
2. To condition (disinfect) your lenses, leave them in a recommended storage solution for at least 4 hours prior to usage or as indicated on the product label.
3. Remove the lens from the case and rinse it with wetting and soaking solution.
4. Place the lens on the tip of your index finger, concave side up.

LENS PLACEMENT

1. Retract the patient's lids with your index finger and thumb.
2. Direct the patient to look straight ahead and place the lens on the cornea.
3. Slowly release the lids and ask the patient to blink. This will center the lens.

LENS REMOVAL

1. Place your index fingers on the lid margins and direct the patient to look straight ahead.
2. Separate the lids, then push them together to remove the lens.

IN-OFFICE CLEANING, DISINFECTION AND STORAGE

FluoroPerm rigid gas permeable contact lenses must be both cleaned and disinfected each time they are removed from the eye. One procedure does not replace the other. Cleaning is necessary to remove mucus and film from the lens surface. Disinfecting is necessary to destroy harmful germs. Leave the FluoroPerm rigid gas permeable contact lenses in a storage solution (such as Unique-pH[®] Multi-Purpose Solution from Alcon Laboratories, Inc.) for a minimum of 4 hours or as indicated on the product label.

To minimize lens warpage during cleaning, the lenses should be cleaned in the palm of the hand rather than between the thumb and fingers.

The directions from any lens care systems used should be followed. Failure to adhere to these procedures may result in the development of serious ocular infections.

Do not mix or alternate the disinfection and storage systems unless so indicated on the product label.

LENS FITTING

General Information

FluoroPerm[®] 92 (paflucocon A), FluoroPerm[®] 60 (paflucocon B) and FluoroPerm[®] 151 (paflucocon D) rigid gas permeable contact lenses may be fitted for daily wear or extended wear using the standard techniques for rigid contact lenses. A diagnostic lens fitting procedure is recommended, although not always required.

FluoroPerm[®] 30 (paflucocon C) rigid gas permeable contact lenses may be fitted for daily wear using the standard techniques for rigid contact lenses. A diagnostic lens fitting procedure is recommended, although not always required.

When placed on the human cornea, the FluoroPerm rigid gas permeable contact lens acts as a refracting medium to focus light rays upon the retina.

Clinical studies have demonstrated that rigid gas permeable contact lenses manufactured from these fluorosilicone acrylate contact lens materials are safe and effective for their intended use. However, the clinical studies may not have included all design configurations or lens parameters that are presently available in these materials. Consequently, when selecting an appropriate lens design and parameters, the eye care practitioner must consider all factors that affect lens performance and ocular health. The potential impact of these factors must be weighted against the patient's needs. Therefore, the continuing ocular health of the patient and lens performance on the eye should be carefully monitored.

Pretrial Examination

A complete contact lens examination should be carried out including general health, previous contact lens history, refraction, keratometry and slit lamp examination. Patients who have evidence of any disease affecting the cornea or conjunctiva, acute or subacute inflammation of the anterior segment of the eye, insufficiency of the lacrimal secretion, corneal hypoesthesia, any disease or infection which will affect the eye or be exacerbated by the wearing of contact lenses are not candidates for wearing these lenses.

FITTING PROCEDURE – SPHERICAL AND ASPHERIC

Selection of Patients

Patients should be individuals who require a daily wear or extended wear lens; are not-aphakic; and, have non-diseased eyes. Patients should have a refractive error which does not exceed 20.00 diopters of myopia and 12.00 diopters of hyperopia for daily wear, or 8.00 diopters of hyperopia for extended wear; and, who may exhibit corneal astigmatism up to 4.00 diopters. Patients predisposed to excessive edema or staining may not be suitable.

Diagnostic Lens Fitting Procedure

Fitting is recommended to be carried out with the aid of diagnostic lenses. Nearly every patient begins with a 9.2 mm trial lens. The exceptions would be patients who have keratometer readings below 41.00 D or above 45.00 D. If the corneal reading is below 41.00 D and the patient appears to have an exceptionally large cornea and palpebral fissure, begin with a 9.6 mm diameter lens. This is rare. Alternatively, if a patient has a keratometer reading which is greater than 45.00 D or has an unusually small palpebral aperture, begin the fitting with an 8.8 mm diameter lens.

The base curve of the lens may be found in Table 1. From the keratometer readings, find the flattest K and steepest K. Enter the table on the left with the corneal cylinder (ΔK) value and follow across to the first diagnostic lens base curve to be used.

TABLE 1

<u>Corneal Cylinder (ΔK)</u>	<u>Lens Base Curve</u>
Plano	0.25 D flatter to on flat K
0.25 – 0.75 D	on flat K to 0.25 D steeper
1.00 – 1.75 D	0.25 D steeper to 0.75 D steeper
>2.00 D	0.75 D steeper to 1.00 D steeper

Place the diagnostic lens on the patient's cornea. Wait approximately 10 minutes for tearing to subside. Check the lens for positioning and movement. Ideally, the lens will ride up with the blink and then quickly drop to a position that is near the center of the cornea. If the lens should drop to the lower limbus or position eccentrically, remove the lens and replace it with a lens that is 0.4 mm larger. If centering is still not achieved, use a diagnostic lens that is 0.25 D steeper.

The power for the final lens may be most accurately determined by an overrefraction with the diagnostic lens in place. This may be carried out with either a trial frame or refractor. Allow approximately 15 minutes for the lens to settle on the patient's eye before performing the overrefraction. Only the spherical power needs to be determined in order to arrive at the final correction.

When the optimum overrefraction has been obtained, the correction should be added to the power of the diagnostic lens to arrive at the final prescription.

EXAMPLE:	Overrefraction	-1.25 D
	Diagnostic Lens	<u>(+)-3.00 D</u>
	Lens Power Ordered	-4.25 D

A patient's lens power requirement may be determined without diagnostic lenses by:

1. Converting the spectacle Rx to minus cylinder form.
2. Adjusting the spectacle Rx for vertex distance.
3. Using the sphere power only.

The selection of the FluoroPerm rigid gas permeable contact lens may be aided by an examination of the fluorescein pattern. The ideal fluorescein pattern shows a definite green central area with some darkening near the periphery of the optic zone. The peripheral curve area should show a definite green band. The most important faulty lens fluorescein pattern to be detected is a corneal seal at the intermediate lens position. The fluorescein pattern shows a dark touch ring that runs completely around the contact lens in the intermediate zone. The larger the lens the greater is the tendency for this to happen. It also occurs most frequently when the optic zone of the contact lens is greater than an 8.0 mm diameter. Hence, a lens design with an optic zone of 8.0 mm or less is usually preferred.

NOTE: Evaluation of fluorescein patterns for lenses manufactured in materials with the UV absorber Uvinul 49 is best accomplished using a slit lamp or Burton Lamp with a Wratten filter over the light source.

A 9.6 mm (large) lens may be needed for patients who have a problem of flare with smaller diameter contact lenses. The flare may occur if the patient has either a very large pupil or if the smaller lens rides off the center of the cornea. Fortunately, flare is only an occasional problem with the 9.6 mm lens because the peripheral curve and edge are specifically designed to minimize the flare problem.

Some patients develop problems of staining, which are due to an optic zone that is too large. A 7.8 mm optic zone is large enough to avoid flare in nearly every contact lens patient. If the patient requires an even larger lens than the 9.2 mm diameter, choose the 9.6 mm diameter lens but keep the optic zone at 8.0 mm in all but the most unusual cases. In this way some tear pumping is achieved in nearly every case and avoids the corneal punctate staining that is due to inadequate lens pumping.

FITTING PROCEDURE – MONOCENTRIC BIFOCAL

Patient Selection

Patients should be selected who require a daily wear lens; are not-aphakic; and, have nondiseased eyes. Patients should have a refractive error which does not exceed 20.00 diopters of myopia and 12.00 diopters of hyperopia; and, who may exhibit corneal astigmatism up to 4.00 diopters and up to +4.00 diopters of add power. Patients predisposed to excessive edema or staining may not be suitable.

Power Compensation

Higher plus power spectacle prescriptions at distance require higher plus distance and add powers, while higher minus power spectacle prescriptions require less minus power and less add powers in bifocal contact lenses.

Prism

The purpose of the prism is to orient the lens, give a low reading position, and provide a supporting edge at the bottom to enable upward displacement by the lower lid.

<u>DISTANT POWER</u>	<u>PRISM</u>
>+8.00	1.00
+4.00	1.25
+2.00	1.50
Plano	1.75
-2.00	2.00
-4.00	2.25
-6.00	2.50
>-8.00	2.75

Seg Height

The average seg height is determined by the diagnostic lens selection and fitting evaluation.

Diagnostic Lens Fitting Procedure

Fitting is recommended to be carried out with the aid of diagnostic lenses. A suggested diagnostic lens set follows.

Base Curves	7.30 to 8.20 mm (.10 mm steps)
Size	9.2 mm
Optic Zone	7.8 mm
Power	- 3.00 diopters
Add	+ 2.00 diopters
Prism Ballast	1.50 diopters
Seg. Height	4.3 mm

The base curve of the lens may be found in Table 2. From the keratometer readings, find the flattest K and steepest K. Find the corneal cylinder (ΔK) value in the left column and follow across, this is the first diagnostic lens base curve to be used.

TABLE 2

<u>Corneal Cylinder (ΔK)</u>	<u>Lens Base Curve</u>
Plano	0.75 D flatter to 0.50 D flatter
0.25 - 0.75 D	0.50 D flatter to 0.25 D flatter
1.00 - 1.75 D	0.25 D flatter to on flat K
>2.00 D	Not more than 0.25 of difference steeper cylinder

Place the diagnostic lens on the patient's cornea. Wait approximately 10 minutes for tearing to subside and check the lens for positioning and movement. Ideally, the lens will ride up with the blink and then quickly drop to a position that is near the center of the cornea. If the lens should drop to the lower limbus or position eccentrically, remove the lens and replace it with a lens that is 0.4 mm larger. If centering is still not achieved use a diagnostic lens which is 0.50 D steeper.

The lens should translate up, with down gaze, so that the segment is within the visual axis for reading. The prism ballast should not rotate greater than 20 degrees nasally, and seg height, in primary gaze, should be even or slightly above lower pupillary margin. If the lens rotates excessively, a larger prism diopter ballast will be necessary.

The power for the final lens may be most accurately determined by an overrefraction with the diagnostic lens in place. This may be carried out with either a trial frame or refractor. Allow approximately 15 minutes for the lens to settle on the patient's eye before performing the overrefraction. Only the spherical power needs to be determined in order to arrive at the final correction.

When the optimum overrefraction has been obtained, the correction should be added to the power of the diagnostic lens to arrive at the final prescription.

EXAMPLE:	Overrefraction	-1.25 D
	Diagnostic Lens	<u> (+)-3.00 D</u>
	Lens Power Ordered	-4.25 D

The selection of the FluoroPerm rigid gas permeable bifocal contact lens may be aided by an examination of the fluorescein pattern. The ideal fluorescein pattern shows a definite green central area with some darkening near the periphery of the optic zone. The peripheral curve area should show a definite green band. The most important faulty lens fluorescein pattern to be detected is a corneal seal at the intermediate lens position. The fluorescein pattern shows a dark touch ring that runs completely around the contact lens in the intermediate zone. The larger the lens the greater is the tendency for this to happen. It also occurs most frequently when the optic zone of the contact lens is greater than an 8.0 mm diameter. Hence, a lens design with an optic zone of 8.0 mm or less is usually preferred.

NOTE: Evaluation of fluorescein patterns for lenses manufactured in materials with the UV absorber Uvinul 49 is best accomplished using a slit lamp or Burton Lamp with a Wratten filter over the light source.

A larger lens may be needed for patients who have a problem of flare with smaller diameter contact lenses. The flare may occur if the patient has either a very large pupil or if the smaller lens rides off the center of the cornea. Fortunately, flare is only an occasional problem with the 9.6 mm lens because the peripheral curve and edge are specifically designed to minimize the flare problem.

Some patients develop problems of staining, which are due to an optic zone that is too large. A 7.8 mm optic zone is large enough to avoid flare in nearly every contact lens patient. If the patient requires an even larger lens than the 9.2 mm diameter, choose the 9.6 mm diameter lens but keep the optic zone at 8.0 mm in all but the most unusual cases. In this way some tear pumping is achieved in nearly every case and avoids the corneal punctate staining that is due to inadequate lens pumping.

FITTING PROCEDURE – CONCENTRIC BIFOCAL

Patient Selection

Patients should be selected who require a daily wear lens; are not-aphakic; and, who have nondiseased eyes. Patients should have a refractive error which does not exceed 20.00 diopters of myopia and 12.00 diopters of hyperopia; and, who may exhibit corneal astigmatism up to 4.00 diopters and up to +4.00 diopters of add power. Patients predisposed to excessive edema or staining may not be suitable.

Power Compensation

Higher plus power spectacle prescriptions at distance require higher plus distance and add powers, while higher minus power spectacle prescriptions require less minus power and less add powers in bifocal contact lenses.

Seg Diameter

The average seg diameter is determined by the diagnostic lens selection and fitting evaluation.

Diagnostic Lens Fitting Procedure

Fitting is recommended to be carried out with the aid of diagnostic lenses. A suggested diagnostic lens set would be as follows.

Base Curves	7.30 to 8.20 mm (.10 mm steps)
Size	9.2 mm
Optic zone	7.8 mm
Power	- 3.00 diopters
Add power	+2.00 diopters
Add diameter	2 to 4 mm

The base curve of the lens may be found in Table 3. From the keratometer readings, find the flattest K and steepest K. Find the corneal cylinder (ΔK) value in the left column and follow across, this is the first diagnostic lens base curve to be used.

TABLE 3

<u>Corneal Cylinder (ΔK)</u>	<u>Lens Base Curve</u>
PLANO	0.25 D flatter to on flat K
0.25 - 0.75 D	On flat K to 0.25 steeper
1.00 - 1.75 D	0.25 D steeper to 0.50 D steeper
2.00 - 2.75 D	0.50 D steeper to 0.75 D steeper
> 2.75 D	0.25 D steeper to 1.00 D steeper

Place the diagnostic lens on the patient's cornea. Wait approximately 10 minutes for tearing to subside and check the lens for positioning and movement. Ideally, the lens will ride up with the blink and then quickly drop to a position which is near the center of the cornea. If the lens should drop to the lower limbus or position eccentrically, remove the lens and replace it with a lens that is 0.4 mm larger. If centering is still not achieved use a diagnostic lens which is 0.25 D steeper.

The power for the final lens may be most accurately determined by an overrefraction with the diagnostic lens in place. This may be carried out with either a diagnostic frame or refractor. Allow approximately 15 minutes for the lens to settle on the patient's eye before performing the overrefraction. Only the spherical power needs to be determined in order to arrive at the final correction. Both distance and near acuity should be checked.

When the optimum overrefraction has been obtained, the correction should be added to the power of the diagnostic lens to arrive at the final prescription.

EXAMPLE:	Overrefraction	-1.25 D
	<u>Diagnostic Lens</u>	<u>(+) -3.00 D</u>
	Lens power ordered	-4.25 D

The selection of the FluoroPerm rigid gas permeable bifocal contact lens may be aided by an examination of the fluorescein pattern. The ideal fluorescein pattern shows a definite green central area with some darkening near the periphery of the optic zone. The peripheral curve area should show a definite green band. The most important faulty lens fluorescein pattern to be detected is a corneal seal at the intermediate lens position. The fluorescein pattern shows a dark touch ring that runs completely around the contact lens in the intermediate

zone. The larger the lens the greater is the tendency for this to happen. It also occurs most frequently when the optic zone of the contact lens is greater than an 8.0 mm diameter. Hence, a lens design with an optic zone of 8.0 mm or less is usually preferred.

NOTE: Evaluation of fluorescein patterns for lenses manufactured in materials with the UV absorber Uvinul 49 is best accomplished using a slit lamp or Burton Lamp with a Wratten filter over the light source.

A larger lens may be needed for patients who have a problem of flare with smaller diameter contact lenses. The flare may occur if the patient has either a very large pupil or if the smaller lens rides off the center of the cornea. Fortunately, flare is only an occasional problem with the 9.60 mm lens because the peripheral curve and edge are specifically designed to minimize the flare problem.

FITTING PROCEDURE – ASPHERIC BIFOCAL

Patient Selection

Patients should be selected who require a daily wear lens; are not-aphakic; and, have nondiseased eyes. Patients should have a refractive error which does not exceed 20.00 diopters of myopia and 12.00 diopters of hyperopia; and, who may exhibit corneal astigmatism up to 4.00 diopters and up to +4.00 diopters of add power. Patients predisposed to excessive edema or staining may not be suitable.

Diagnostic Lens Fitting Procedure

Fitting is recommended to be carried out with the aid of diagnostic lenses. A suggested diagnostic lens set follows.

Base Curves	7.30 to 8.20 mm (.10 mm steps)
Size	9.2 mm
Optic Cone	7.8 mm
Power	- 3.00 diopters
Add	+2.00 diopters

The base curve of the diagnostic lens should be 2.75 diopters steeper than the flattest keratometer reading.

EXAMPLE:	K Reading	42.50 x 43.50
	<u>+2.75 Steep</u>	<u>2.75</u>
	Diagnostic Lens Base Curve	45.25 = 7.456 mm

Place the diagnostic lens on the patient's cornea. Wait approximately 10 minutes for tearing to subside and check the lens for positioning and movement. Ideally, the lens will ride up with the blink and then quickly drop to a position that is near the center of the cornea. If the lens should drop to the lower limbus or position eccentrically, remove the lens and replace it with a lens that is 0.4 mm larger. If centering is still not achieved, use a diagnostic lens that is 0.50 D steeper.

The power for the final lens may be most accurately determined by an overrefraction with the diagnostic lens in place. This may be carried out with either a diagnostic frame or refractor. Allow approximately 15 minutes for the lens to settle on the patient's eye before performing the overrefraction. Only the spherical power needs to be determined in order to arrive at the final correction. Both distance and near acuity should be checked.

With the diagnostic lenses on the eyes, place the overrefraction in a diagnostic frame for both eyes. (The use of a phoropter is not recommended for this test.)

- Add a +0.50 sphere to both sides of trial frame and ask the patient to read the near point card. Then ask the patient to read the 20/20 line on the distance chart (not how clearly he can read).
- Continue adding +0.25 at a time until the patient cannot read all of the 20/20 line.
- If the patient can read most of the 20/20 line, and at least J-3 on the near point card, the total power of the contact lens power and the power of diagnostic lenses from the trial frame should be ordered.

When the optimal overrefraction has been obtained, the correction should be added to the power of the diagnostic lens to arrive at the final prescription.

EXAMPLE:	Overrefraction	-1.25 D
	<u>Diagnostic Lens</u>	<u>(+) -3.00 D</u>
	Lens power ordered	-4.25 D

The selection of the FluoroPerm rigid gas permeable bifocal contact lens may be aided by an examination of the fluorescein pattern. The ideal fluorescein pattern shows a definite green central area with some darkening near the periphery of the optic zone. The peripheral curve area should show a definite green band. The most important faulty lens fluorescein pattern to be detected is a corneal seal at the intermediate lens position. The fluorescein pattern shows a dark touch ring that runs completely around the contact lens in the intermediate zone. The larger the lens the greater is the tendency for this to happen. It also occurs most frequently when the optic zone of the contact lens is greater than an 8.0mm diameter. Hence, a lens design with an optic zone of 8.0mm or less is usually preferred.

NOTE: Evaluation of fluorescein patterns for lenses manufactured in materials with the UV absorber Uvinul 49 is best accomplished using a slit lamp or Burton Lamp with a Wratten filter over the light source.

A larger lens may be needed for patients who have a problem of flare with smaller diameter contact lenses. The flare may occur if the patient has either a very large pupil or if the smaller lens rides off the center of the cornea. Fortunately, flare is only an occasional problem with the 9.60 mm lens because the peripheral curve and edge are specifically designed to minimize the flare problem.

FITTING PROCEDURE – TORIC

Patient Selection

Patients should be selected who require a daily wear lens; are not-aphakic; and, who have nondiseased eyes. Patients should have a refractive error that does not exceed 20.00 diopters of myopia and 12.00 diopters of hyperopia; and, who may exhibit corneal astigmatism up to 6.00 diopters.

Diagnostic Lens Fitting Procedure

The base curve of the lens may be found in Table 4. From the keratometer readings, find the flattest K and steepest K. Find the corneal cylinder (ΔK) value in the left column and follow across, this is the first diagnostic set to be used.

TABLE 4

<u>Corneal Cylinder (ΔK)</u>	<u>Diagnostic Set</u>
1.00 - 2.50	2 Diopter Diagnostic Set
2.75 - 4.00	3 Diopter Diagnostic Set
4.25 – Up	4 Diopter Diagnostic Set

Place the diagnostic lens on the patient's cornea. Wait approximately 10 minutes for tearing to subside and check the lens for positioning and movement. Ideally, the lens will ride up with the blink and then quickly drop to a position that is near the center of the cornea. If the lens should drop to the lower limbus or position eccentrically, remove the lens and replace it with a lens that is 0.4 mm larger. If centering is still not achieved, use a diagnostic lens that is 0.50 D steeper.

Other Diagnostic Lens Procedure

Select a bitoric diagnostic lens that provides alignment bearing and positions the lens slightly beneath the upper lid.

Allow approximately 15 minutes for the lens to settle on the patient's eye before performing the overrefractions.

Perform a sphero-cylinder overrefraction, adding the sphere power from the overrefraction to the flattest meridian of the diagnostic lens.

EXAMPLE:	Diagnostic Lens	42.00	x	44.00
		-1.00		-2.00
	Overrefraction	<u>-2.00</u>		
	Order	-3.00		-2.00

If the cylinder finding is -0.50 or less, order cylinder power of diagnostic lens.

EXAMPLE:	Diagnostic Lens	42.00	x	44.00
		-1.00		-2.00
	Overrefraction	<u>PL</u>		<u>-0.50</u>
	Order	-1.00		-2.00

If cylinder finding is -0.75 or larger, and the axis is at or near the diagnostic lens axis, add the minus cylinder power to the steep meridian of the diagnostic lens.

EXAMPLE:	Diagnostic Lens	42.00	x	44.00
		-1.00		-2.00
	Overrefraction	<u>PL</u>		<u>-1.00</u> x 180
	Order	-1.00		-3.00

The selection of the FluoroPerm rigid gas permeable toric contact lens may be aided by an examination of the fluorescein pattern. The ideal fluorescein pattern shows a definite green central area with some darkening near the periphery of the optic zone. The peripheral curve area should show a definite green band.

NOTE: Evaluation of fluorescein patterns for lenses manufactured in materials with the UV absorber Uvinul 49 is best accomplished using a slit lamp or Burton Lamp with a Wratten filter over the light source.

A larger lens may be needed for patients who have a problem of flare with smaller diameter contact lenses. The flare may occur if the patient has either a very large pupil or if the smaller lens rides off the center of the cornea. Fortunately, flare is only an occasional problem.

FITTING PROCEDURE – KERATOCONUS

Patient Selection

Keratoconus is a non-inflammatory ocular condition in which the cornea progressively thins causing a cone-like bulge to develop. As the cornea steepens the anterior corneal surface (epithelium) becomes irregular resulting in visual impairment. This irregularity cannot be completely corrected with spectacles – instead a RGP contact lens* is used, becoming the new anterior refracting surface. FluoroPerm® contact lenses for keratoconus are indicated for patients who have a demonstrated need as described, and a desire for refractive correction with RGP contact lenses and who do not have any of the contraindications for RGP contact lenses. Refer the Package Insert, Contraindications (Reasons Not To Use).

Special Fitting Considerations

FluoroPerm® contact lenses for keratoconus are designed to be fitted so as to optically correct irregular astigmatism and thereby improve visual acuity. The lens design and the manner in which the lens is fitted are intended to work together to accomplish this goal. The keratoconus design utilizes smaller optic zone diameters, steeper base curves, spherical and/or aspherical periphery curves to closely approximate the unusual topography typical in patients with keratoconus. For example, keratoconus lens designs such as the McGuire Cones utilize small posterior optic zones and a series of peripheral curves to achieve this fitting relationship.

*See “NOTE” on page 15.

FluoroPerm[®] contact lenses for keratoconus may be fitted using a modification of the standard techniques for rigid gas permeable contact lenses.

Extended wear lenses should not be used to correct keratoconus.

Initial Lens Power Selection

Standard procedures for determining power of rigid gas permeable contact lenses may be used, including compensation for vertex distance.

Initial Lens Diameter Selection

Usually, lens diameters from 7.0 mm to 11.5 mm are chosen to maximize positioning on the cornea and to minimize lens movement. Lens diameters outside of this range are occasionally used. This guide is only a general recommendation and the specification for an individual patient will depend on the eye care practitioner's professional judgment.

Lens diameter is primarily a function of the base curve but may be influenced by power (plus lenses require a larger diameter to compensate for weight) and anatomical considerations (small palpebral opening, excessively large pupil, etc.) and the patient's corneal topography.

Initial Lens Base Curve Selection

The base curve of the first lens fitted is generally equal to or slightly steeper than the flattest keratometry reading to achieve an apical clearance or apical alignment fitting relationship.

Initial Lens Evaluation

Blink induced lens movement should show downward lens movement with the lid motion (average 1 mm) and then upward with the lid motion (average 1 mm) as with a standard RGP contact lens. During the interblink period the lens should have little or no motion (average less than 1 mm).

The lens should position centrally or slightly inferiorly, as it will tend to migrate to the steepest corneal area.

A lens that is too tight will show reduced movement upon blinking. Bubbles may be detected behind the lens.

A lens that is too loose will move excessively on the cornea following each blink. The lens may ride in either a position that is too high or too low or in an eccentric position. A loose lens is usually uncomfortable for the patient.

Diagnostic Lens Fitting Procedure

Diagnostic (trial) lens fitting is recommended whenever possible. Trial lens fitting allows a more accurate determination of lens specifications for the lens fit and power. Choose the first lens according to the base curve selection criteria for the specific lens design. Trial lenses are essential in fitting patients whose corneal topography is distorted.

Select a trial lens and place the lens on the eye. Evaluate the lens using white light for the following:

Centering

Lenses may not center well due to the unusual corneal topography in patients with keratoconus. Often the lens will position inferiorly over the steepest corneal area.

Movement

Lens movement should be equivalent to or slightly less than a standard RGP lens.

Fluorescein Pattern

The fluorescein pattern should show a lens with either mild apical clearance or “feather touch” (alignment) over the steepest conical area. In the periphery there should be another area of alignment and near the edge a thin band of pooling.

The fluorescein pattern provides the best method for monitoring the contact lens fit over time.

Special Follow-up Care

With lenses in place on the eyes, evaluate fitting performance to assure that the criteria of a well-fitted lens continue to be satisfied. The fluorescein pattern provides a guide to lens adaptation. If the lens demonstrates reduced movement consider exchanging for another of flatter base curve. Usually, a lens with a 0.50 diopters flatter base curve should be the next choice with variations from this based on the judgment of the eye care practitioner.

A lens with excessive movement should be replaced with another that is 0.50 diopters steeper base curve.

After lens removal, conduct a thorough biomicroscopy examination to detect the presence of unusual vertical corneal striae in the posterior central cornea and/or corneal neovascularization. Note: some vertical striae are typical in advanced stages of keratoconus. The presence of these conditions may be indicative of excessive corneal edema.

The recommended schedule for follow-up visits is the same as standard lenses. Reference “Follow-up Patient Care”, page 19.

NOTE: Practitioners should consult their finishing lab for available keratoconus lens designs. The design parameters must meet the parameters specified in the product labeling.

MANDELL-MOORE BITORIC LENS GUIDE

Contact lens practitioners can now incorporate the FluoroPerm material advantages in lens designs for their astigmatic patients. The following fitting procedure and charts are derived from Mandell and Moore.¹

As with spherical FluoroPerm lenses, you must first evaluate for contraindications to lens wear. Very precise keratometry and refraction will be necessary. You will need to determine the flat K and steep K readings and record the refraction in minus cylinder form.

The overall and optic zone diameters are chosen using the same criteria as with spherical lens designs. Generally, bitoric lens diameters are about 0.2 mm smaller than spherical lenses designed for intrapalpebral fitting. It is important to avoid lid attachment since lid action may cause the lens to rotate from the intended axis.

There are two opposing considerations when selecting the toric posterior surface of a bitoric contact lens in order to achieve the optimum fit. The toric surface must conform close enough to the corneal contour to minimize rotation of the lens. However, some deviation from perfect lens-to-cornea conformation is needed to create pumping of the tear fluid.

The base curve in the flatter meridian should usually be made 0.25 D flatter than the cornea. The base curve in the steeper meridian should be 0.50 to 1.25 D flatter than the cornea depending on the amount of corneal astigmatism. This additional "fit factor" is summarized in the following table.

¹ Mandell, R.G., Moore, C.F.: A Bitoric Lens Guide That Really Is Simple. Contact Lens Spectrum, November, 1988, 83-85.

Right Eye

KERATOMETRY 44.00 @ 180 47.25 @ 90

SPECTACLE Rx (MINUS CYL FORM) -3.50 - 3.50 x 180

	FLATTEST K	SPHERE POWER	STEEPEST K	SPH +CYL POWER
1. ENTER K	44.00	XXXXXXXXXXXXXX	47.25	XXXXXXXXXXXXXX
2. ENTER SPECTACLE POWER		-3.50		-7.00
3. VERTEX CORRECTED		-3.50		-6.50
4. FIT FACTORS	(-) 0.25	(+) 0.25	(-) 0.75	(+) 0.75
ADD LINES	1&4	3&4	1&4	3&4
5. FINAL C.L. Rx	43.75	-3.25	46.50	-5.75
	BASECURVE	POWER	BASECURVE	POWER

Left Eye

KERATOMETRY 42.00 @ 180 46.50 @ 90

SPECTACLE Rx (MINUS CYL FORM) +12.50 - 4.00 x 180

	FLATTEST K	SPHERE POWER	STEEPEST K	SPH +CYL POWER
1. ENTER K	42.00	XXXXXXXXXXXXXX	46.50	XXXXXXXXXXXXXX
2. ENTER SPECTACLE POWER		+12.50		8.50
3. VERTEX CORRECTED		+14.75		9.50
4. FIT FACTORS	(-) 0.25	(+) 0.25	(-) 1.00	(+) 1.00
ADD LINES	1&4	3&4	1&4	3&4
5. FINAL C.L. Rx	41.75	+15.00	45.50	+10.50
	BASECURVE	POWER	BASECURVE	POWER

VERTEX DISTANCE CORRECTION					
4.00	3.75	8.00	7.25	12.00	10.50
4.25	4.00	8.25	7.50	12.25	10.75
4.50	4.25	8.50	7.75	12.50	10.75
4.75	4.50	8.75	8.00	12.75	11.00
5.00	4.75	9.00	8.00	13.00	11.25
5.25	5.00	9.25	8.25	13.25	11.25
5.50	5.25	9.50	8.50	13.50	11.50
5.75	5.50	9.75	8.75	13.75	11.75
6.00	5.50	10.00	9.00	14.00	12.00
6.25	5.75	10.25	9.00	14.25	12.00
6.50	6.00	10.50	9.25	14.50	12.25
6.75	6.25	10.75	9.50	14.75	12.50
7.00	6.50	11.00	9.75	15.00	12.50
7.25	6.75	11.25	10.00	15.25	12.75
7.50	7.00	11.50	10.00	15.50	13.00
7.75	7.00	11.75	10.25	15.75	13.00

CORNEAL CYL, D	FIT FACTOR	
	FLAT MERIDIAN	STEEP MERIDIAN
2.00	ON K	.50 FLATTER
2.50	.25 FLATTER	.50 "
3.00	.25 "	.75 "
3.50	.25 "	.75 "
4.00	.25 "	1.00 "
5.00	.25 "	1.25 "

If the spectacle lens power is less than 4.00 diopters then line 3 = line 2. Otherwise: For minus power spectacle lenses find the power in the left side of the column and convert to the power in the right side, but retain the minus sign. For plus power spectacle lenses find the power in the right side of the column and convert to the power in the left side, but retain the plus sign.

Right Eye

KERATOMETRY	@	@	
SPECTACLE Rx (MINUS CYL FORM)	X		

1. ENTER K 2. ENTER SPECTACLE POWER 3. VERTEX CORRECTED 4. FIT FACTORS ADD LINES 5. FINAL C.L. Rx	FLATTEST K	SPHERE POWER	STEEPEST K	SPH +CYL POWER	
		XXXXXXXXXXXXXXXXXX		XXXXXXXXXXXXXXXXXX	
		(-)	(+)	(-)	(+)
		1 & 4	3 & 4	1 & 4	3 & 4
	BASECURVE	POWER	BASECURVE	POWER	

Left Eye

KERATOMETRY	@	@	
SPECTACLE Rx (MINUS CYL FORM)	X		

1. ENTER K 2. ENTER SPECTACLE POWER 3. VERTEX CORRECTED 4. FIT FACTORS ADD LINES 5. FINAL C.L. Rx	FLATTEST K	SPHERE POWER	STEEPEST K	SPH +CYL POWER	
		XXXXXXXXXXXXXXXXXX		XXXXXXXXXXXXXXXXXX	
		(-)	(+)	(-)	(+)
		1 & 4	3 & 4	1 & 4	3 & 4
	BASECURVE	POWER	BASECURVE	POWER	

VERTEX DISTANCE CORRECTION					
4.00	3.75	8.00	7.25	12.00	10.50
4.25	4.00	8.25	7.50	12.25	10.75
4.50	4.25	8.50	7.75	12.50	10.75
4.75	4.50	8.75	8.00	12.75	11.00
5.00	4.75	9.00	8.00	13.00	11.25
5.25	5.00	9.25	8.25	13.25	11.25
5.50	5.25	9.50	8.50	13.50	11.50
5.75	5.50	9.75	8.75	13.75	11.75
6.00	5.50	10.00	9.00	14.00	12.00
6.25	5.75	10.25	9.00	14.25	12.00
6.50	6.00	10.50	9.25	14.50	12.25
6.75	6.25	10.75	9.50	14.75	12.50
7.00	6.50	11.00	9.75	15.00	12.50
7.25	6.75	11.25	10.00	15.25	12.75
7.50	7.00	11.50	10.00	15.50	13.00
7.75	7.00	11.75	10.25	15.75	13.00
				20.00	16.00

CORNEAL CYL, D	FIT FACTOR	
	FLAT MERIDIAN	STEEP MERIDIAN
2.00	ON K	.50 FLATTER
2.50	.25 FLATTER	.50 "
3.00	.25 "	.75 "
3.50	.25 "	.75 "
4.00	.25 "	1.00 "
5.00	.25 "	1.25 "

If the spectacle lens power is less than 4.00 diopters then line 3 = line 2. Otherwise: For minus power spectacle lenses find the power in the left side of the column and convert to the power in the right side, but retain the minus sign. For plus power spectacle lenses find the power in the right side of the column and convert to the power in the left side, but retain the plus sign.

THE CLINICAL PICTURE

With the ideal fit the lens should move freely with the lid during a blink and then drop quickly to a position near the center of the cornea. In some patients, the lens will ride slightly high. This is most desirable. It is especially favorable if the lens rides slightly under the upper lid since that will reduce lens edge sensation and make the lens most comfortable. It is best to avoid having the lens ride exceptionally high so that excessive lid pressure is exerted on the superior lens margin. Over an extended wearing period, this inevitably leads to structural changes in the superior corneal epithelium. If the lens appears to center well and move adequately following the blink, proceed to determine the refractive correction.

A lens that is too tight will show reduced movement upon blinking. The lens usually occupies a centered corneal position and may not move far from this position. Bubbles may be detected in the post-lens space.

A lens that is too loose will move excessively on the cornea following each blink. The lens may ride in either a position that is too high, too low or in an eccentric position. A loose lens is usually uncomfortable for the patient.

FOLLOW-UP PATIENT CARE

Follow-up examination should include an evaluation of lens movement, centration, comfort and fluorescein pattern. Lens movement will decrease as the tear volume is diminishing during adaptation. An assessment of vision and eye health, including inspection of the cornea for edema and/or staining, should be performed. Patient symptoms should also be assessed.

RECOMMENDED DAILY WEAR FOLLOW-UP SCHEDULE

First Follow-up Examination -	Immediately following 3 hours of lens wear on the 1 st , 2 nd , or 3 rd day following dispensing
Second Follow-up Examination -	After 1 week of lens wear
Third Follow-up Examination -	After 3 weeks of lens wear
Subsequent Follow-up Examinations -	After 2 months of lens wear; then regular check-ups as determined by the eye care practitioner

RECOMMENDED EXTENDED WEAR FOLLOW-UP SCHEDULE

First Follow-up Examination -	Immediately following the first overnight wearing of the lens
Second Follow-up Examination -	After 3 days of extended wear of the lens
Third Follow-up Examination -	After 1 week of extended wear of the lens
Fourth Follow-up Examination -	After 1 month of extended wear of the lens
Subsequent Follow-up Examinations -	After 3 and 6 months of extended wear of the lens; then regular check-ups as determined by the eye care practitioner

NOTE: See Package Insert for additional safety information.

WEARING SCHEDULE

See Package Insert (Wearing Schedule) for the maximum suggested wearing time for each product.

PATIENT LENS CARE DIRECTIONS

See Package Insert (Lens Care Directions) for these directions.

CARE FOR A STICKING (nonmoving) LENS

See Package Insert (Care For A Sticking Lens) for these instructions.

HOW SUPPLIED

Each FluoroPerm[®] lens is supplied nonsterile in an individual plastic case. The lens is shipped dry; or, wet shipped in Unique-pH[®] Multi-Purpose Solution. This solution contains hydroxypropyl guar, a unique wetting and conditioning polymer system, polyethylene glycol, Tetronic*, boric acid, propylene glycol; and, is preserved with POLYQUAD[®] (polyquaternium-1) 0.0011% and edetate disodium 0.01%. The case, packing slip or invoice is marked with the base curve radius, dioptric power, diameter, center thickness, inclusion of UV absorber, lot number, fill date and the color of the lens.

Unique-pH[®] is a Trademark of Alcon Laboratories, Inc.
* Registered Trademark of BASF corp.

Never reuse the solution. You may store the lens in the unopened container until ready to dispense, up to a maximum of thirty (30) days from the Fill Date. When a lens has been stored for 30 days in its original packaging solution, it should be cleaned and disinfected with an approved product (see recommended product list in the Lens Care Directions section), and placed into inventory as you presently do with any other RGP lens held in your office. Follow the directions on the selected disinfecting solution regarding prolonged storage.

REPORTING OF ADVERSE REACTIONS

All serious adverse experiences and adverse reactions observed in patients wearing or experienced with the lenses should be reported to the manufacturer.

Paragon Vision Sciences
947 E. Impala Avenue
Mesa, Arizona 85204-6619

1-800-528-8279
1-480-892-7602
1-480-926-7369 FAX

PACKAGE INSERT

FluoroPerm[®] 92 (paflucocon A)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear

FluoroPerm[®] 60 (paflucocon B)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear

FluoroPerm[®] 30 (paflucocon C)

Rigid Gas Permeable Contact Lenses for Daily Wear

FluoroPerm[®] 151 (paflucocon D)

Rigid Gas Permeable Contact Lenses for Daily and Extended Wear

Shipped Dry; or, Wet Shipped in Unique-pH[®] Multi-Purpose Solution

IMPORTANT: Please read carefully and keep this information for future use.

PROBLEMS WITH CONTACT LENSES AND LENS CARE PRODUCTS COULD RESULT IN SERIOUS INJURY TO THE EYE. It is essential that you follow your eye care practitioner's directions and all labeling instructions for proper use of your contact lenses and lens care products. **EYE PROBLEMS, INCLUDING CORNEAL ULCERS, CAN DEVELOP RAPIDLY AND LEAD TO LOSS OF VISION; THEREFORE, IF YOU EXPERIENCE EYE DISCOMFORT, EXCESSIVE TEARING, VISION CHANGES, OR REDNESS OF THE EYE, OR OTHER PROBLEMS WITH YOUR EYES, IMMEDIATELY REMOVE YOUR LENSES, AND PROMPTLY CONTACT YOUR EYE CARE PRACTITIONER.**

FLUOROPERM[®] 92 (paflucocon A)

RIGID GAS PERMEABLE CONTACT LENSES FOR DAILY WEAR AND EXTENDED WEAR

Daily or Extended Wear Spherical, Aspheric contact lenses for:
Nearsightedness (myopia).
Farsightedness (hyperopia).

Daily Wear Bifocal, Toric contact lenses for:
Nearsightedness (myopia).
Farsightedness (hyperopia).

DESCRIPTION

FluoroPerm[®] 92 (paflucocon A) rigid gas permeable contact lenses for daily wear are available as lathe cut or molded firm contact lenses with spherical, aspheric, bifocal or toric anterior and/or posterior; or, bitoric surfaces in clear and tinted versions. FluoroPerm 92 rigid gas permeable contact lenses for extended wear are available as lathe cut or molded firm contact lenses with spherical or aspheric anterior or posterior surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 92 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

FluoroPerm 92 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM[®] 92 (paflucocon A)

Refractive Index	1.453(Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	93%
Luminous Transmittance (Green)	95%
Wetting Angle (Receding Angle) ⁺⁺	16°
Wetting Angle (Receding Angle) ⁺⁺⁺	64°
Specific Gravity	1.10
Hardness (Shore D)	81
Water Content	< 1%
Oxygen Permeability*	92 x 10 ⁻¹¹ Dk at 35° C
Oxygen Permeability**	64 x 10 ⁻¹¹ Dk at 35° C

⁺ Determination of the Spectral and Luminous Transmittance, ISO 8599:1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺ Sessile Drop Technique per ANSI Z80.20. 8.11

* (cm²/sec) (mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec) (mL O₂) / (mL x mm Hg) ISO/ANSI Fatt Method

Lens Parameters:

Chord Diameter	7.0 to 11.5 mm
Center Thickness	0.05 to 0.70 mm
Base Curve	6.50 to 9.00 mm

Powers Daily Wear	-20.00 to +12.00 Diopters
Powers Extended Wear	-20.00 to +8.00 Diopters
Bifocal Add Powers	+0.25 to +4.00 Diopters
Monocentric Bifocal Add Diameter	4.0 to 9.0 mm
Monocentric Bifocal Prism	1.0 to 2.5 Diopters
Concentric Bifocal Add Diameter	2.0 to 4.0 mm

ACTION

FluoroPerm 92 rigid gas permeable toric and bifocal contact lenses are intended for daily wear only. FluoroPerm 92 spherical and aspheric contact lenses are intended for daily or extended wear.

When placed on the human cornea, the FluoroPerm 92 rigid gas permeable contact lens acts as a refracting medium to focus light rays upon the retina.

The toric lens provides for the individual meridional power requirements of the astigmatic eye. In the bifocal lens, the distance or near power prescription is provided in a small area with the near or distance prescription surrounding it.

INDICATIONS (USES)

FluoroPerm[®] 92 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm[®] 92 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm[®] 92 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 92 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm[®] 92 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 92 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

CONTRAINDICATIONS (REASONS NOT TO USE)

FluoroPerm 92 rigid gas permeable contact lenses are contraindicated by the presence of any of the following conditions:

- Acute or subacute inflammations of the anterior segment of the eye.
- Any eye disease, injury, or abnormality, other than keratoconus, that affects the cornea, conjunctive or eyelids.
- Insufficiency of lacrimal secretion (dry eyes).
- Corneal hypoesthesia (reduced corneal sensitivity).
- Any systemic disease, which may affect the eye or be exacerbated by wearing contact lenses.
- Allergic reactions of ocular surfaces or adnexa which may be induced or exaggerated by wearing contact Lenses and/or using contact lens solutions.
- Any active corneal infection (bacterial, fungal or viral).

WEARING SCHEDULE

THE EYE CARE PRACTITIONER SHOULD DETERMINE THE WEARING SCHEDULE. Patients tend to over wear the lenses initially. It is important to adhere to the initial maximum wearing schedule. Regular checkups, as determined by the eye care practitioner, are also extremely important.

The maximum suggested wearing time for FluoroPerm 92 rigid gas permeable contact lenses is:

DAILY WEAR (less than 24 hours while awake)

DAY	1	2	3	4	5	6	7	8	9	10 - 14	15 & after
SUGGESTED HOURS	3	4	5	6	7	8	9	10	11	12	All waking hours
HOURS WORN											

EXTENDED WEAR

Not every patient is able to wear the FluoroPerm 92 rigid gas permeable contact lens on an extended wear basis, even if able to wear the lens for daily wear. An initial 14-day daily wear period is recommended prior to overnight wear. Regular check-ups, as determined by the eye care practitioner, are extremely important. The lens should be removed at least once every week (7 days) for cleaning and disinfection.

Extended wear lenses should not be used to correct keratoconus.

DAY	SUGGESTED HOURS	HOURS WORN
15 (and after)	24 hours	

WARNING

The risk of ulcerative keratitis has been shown to be greater among wearers of extended wear lenses than among wearers of daily wear lenses. The risk among extended wear lens wearers increases with the number of consecutive days that lenses are worn between removals, beginning with the first overnight use. This risk can be reduced by carefully following directions for routine lens care, including cleaning of the lens storage case. Additionally, smoking increases the risk of ulcerative keratitis for contact lens wearers.

It is recommended that contact lens wearers see their eye care practitioners twice each year or, if directed, more frequently.

FLUOROPERM® 60
(paflucocon B)

RIGID GAS PERMEABLE CONTACT LENSES FOR
DAILY WEAR AND EXTENDED WEAR

Daily or Extended Wear Spherical, Aspheric contact lenses for:
Nearsightedness (myopia).
Farsightedness (hyperopia).

Daily Wear Bifocal, Toric contact lenses for:
Nearsightedness (myopia).
Farsightedness (hyperopia).

DESCRIPTION

FluoroPerm® 60 (paflucocon B) rigid gas permeable contact lenses for daily wear are available as lathe cut or molded firm contact lenses with spherical, aspheric, bifocal or toric anterior and/or posterior; or, bitoric

surfaces in clear and tinted versions. FluoroPerm 60 rigid gas permeable contact lenses for extended wear are available as lathe cut or molded firm contact lenses with spherical or aspheric anterior or posterior surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 60 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

FluoroPerm 60 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM® 60 (paflucocon B)

Refractive Index	1.453(Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	95%
Luminous Transmittance (Crystal Blue)	98%
Luminous Transmittance (Green)	95%
Luminous Transmittance (Brown)	85%
Wetting Angle (Receding Angle) ⁺⁺	14.7°
Wetting Angle (Receding Angle) ⁺⁺⁺	62°
Specific Gravity	1.15
Hardness (Shore D)	83
Water Content	< 1%
Oxygen Permeability*	60 x 10 ⁻¹¹ Dk at 35° C
Oxygen Permeability**	43 x 10 ⁻¹¹ Dk at 35° C

⁺ Determination of the Spectral and Luminous Transmittance, ISO 8599:1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺ Sessile Drop Technique per ANSI Z80.20. 8.11

* (cm²/sec) (mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec) (mL O₂) / (mL x mm Hg) ISO/ANSI Fatt Method

Lens Parameters:

Chord Diameter	7.0 to 11.5 mm
Center Thickness	0.05 to 0.70 mm
Base Curve	6.50 to 9.00 mm
Powers Daily Wear	-20.00 to +12.00 Diopters
Powers Extended Wear	-20.00 to +8.00 Diopters
Bifocal Add Powers	+0.25 to +4.00 Diopters
Monocentric Bifocal Add Diameter	4.0 to 9.0 mm
Monocentric Bifocal Prism	1.0 to 2.5 Diopters
Concentric Bifocal Add Diameter	2.0 to 4.0 mm

ACTION

FluoroPerm 60 rigid gas permeable toric and bifocal contact lenses are intended for daily wear only. FluoroPerm 60 spherical and aspheric contact lenses are intended for daily or extended wear.

When placed on the human cornea, the FluoroPerm 60 rigid gas permeable contact lens acts as a refracting medium to focus light rays upon the retina.

The toric lens provides for the individual meridional power requirements of the astigmatic eye. In the bifocal lens, the distance or near power prescription is provided in a small area with the near or distance prescription surrounding it.

INDICATIONS (USES)

FluoroPerm® 60 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm® 60 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm® 60 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm® 60 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm® 60 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm® 60 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

CONTRAINDICATIONS (REASONS NOT TO USE)

FluoroPerm 60 rigid gas permeable contact lenses are contraindicated by the presence of any of the following conditions:

- Acute or subacute inflammations of the anterior segment of the eye.
- Any eye disease, injury, or abnormality, other than keratoconus, that affects the cornea, conjunctive or eyelids.
- Insufficiency of lacrimal secretion (dry eyes).
- Corneal hypoesthesia (reduced corneal sensitivity).
- Any systemic disease, which may affect the eye or be exacerbated by wearing contact lenses.
- Allergic reactions of ocular surfaces or adnexa which may be induced or exaggerated by wearing contact Lenses and/or using contact lens solutions.
- Any active corneal infection (bacterial, fungal or viral).

WEARING SCHEDULE

THE EYE CARE PRACTITIONER SHOULD DETERMINE THE WEARING SCHEDULE. Patients tend to over wear the lenses initially. It is important to adhere to the initial maximum wearing schedule. Regular checkups, as determined by the eye care practitioner, are also extremely important.

The maximum suggested wearing time for FluoroPerm 60 rigid gas permeable contact lenses is:

DAILY WEAR (less than 24 hours while awake)

DAY	1	2	3	4	5	6	7	8	9	10 - 14	15 & after
SUGGESTED HOURS	3	4	5	6	7	8	9	10	11	12	All waking hours
HOURS WORN											

EXTENDED WEAR

Not every patient is able to wear the FluoroPerm 60 rigid gas permeable contact lens on an extended wear basis, even if able to wear the lens for daily wear. An initial 14-day daily wear period is recommended prior to

overnight wear. Regular check-ups, as determined by the eye care practitioner, are extremely important. The lens should be removed at least once every week (7 days) for cleaning and disinfection.

Extended wear lenses should not be used to correct keratoconus.

DAY	SUGGESTED HOURS	HOURS WORN
15 (and after)	24 hours	

WARNING

The risk of ulcerative keratitis has been shown to be greater among wearers of extended wear lenses than among wearers of daily wear lenses. The risk among extended wear lens wearers increases with the number of consecutive days that lenses are worn between removals, beginning with the first overnight use. This risk can be reduced by carefully following directions for routine lens care, including cleaning of the lens storage case. Additionally, smoking increases the risk of ulcerative keratitis for contact lens wearers.

It is recommended that contact lens wearers see their eye care practitioners twice each year or, if directed, more frequently.

FLUOROPERM® 30
(pafluocon C)

RIGID GAS PERMEABLE CONTACT LENSES FOR DAILY WEAR

Spherical, Aspheric, Bifocal, Toric contact lenses for:
 Nearsightedness (myopia).
 Farsightedness (hyperopia).

DESCRIPTION

FluoroPerm® 30 (pafluocon C) rigid gas permeable contact lenses for daily wear are available as lathe cut or molded firm contact lenses with spherical, aspheric, bifocal or toric anterior and/or posterior; or, bitoric surfaces in clear and tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 30 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

FluoroPerm 30 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM® 30 (pafluocon C)

Refractive Index	1.466(Nd at 25°C)
Luminous Transmittance ⁺ (Clear)	99%
Luminous Transmittance (Blue)	94%
Luminous Transmittance (Crystal Blue)	98%
Luminous Transmittance (Majestic Blue)	79%
Luminous Transmittance (Green)	95%
Luminous Transmittance (Gray)	91%

Wetting Angle (Receding Angle) ⁺⁺	12.8°
Wetting Angle (Receding Angle) ⁺⁺⁺	61°
Specific Gravity	1.14
Hardness (Shore D)	84
Water Content	< 1%
Oxygen Permeability*	30 x 10 ⁻¹¹ Dk at 35° C
Oxygen Permeability**	30 x 10 ⁻¹¹ Dk at 35° C

⁺ Determination of the Spectral and Luminous Transmittance, ISO 8599:1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺ Sessile Drop Technique per ANSI Z80.20. 8.11

* (cm²/sec) (mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec) (mL O₂) / (mL x mm Hg) ISO/ANSI Fatt Method

Lens Parameters:

Chord Diameter	7.0 to 11.5 mm
Center Thickness	0.05 to 0.70 mm
Base Curve	6.50 to 9.00 mm
Powers Daily Wear	-20.00 to +12.00 Diopters
Bifocal Add Powers	+0.25 to +4.00 Diopters
Monocentric Bifocal Add Diameter	4.0 to 9.0 mm
Monocentric Bifocal Prism	1.0 to 2.5 Diopters
Concentric Bifocal Add Diameter	2.0 to 4.0 mm

ACTION

FluoroPerm 30 rigid gas permeable spherical, aspheric, toric and bifocal contact lenses are intended for daily wear only.

When placed on the human cornea, the FluoroPerm 30 rigid gas permeable contact lens acts as a refracting medium to focus light rays upon the retina.

The toric lens provides for the individual meridional power requirements of the astigmatic eye. In the bifocal lens, the distance or near power prescription is provided in a small area with the near or distance prescription surrounding it.

INDICATIONS (USES)

FluoroPerm[®] 30 rigid gas permeable contact lenses are indicated for daily wear as recommended by the eye care practitioner.

FluoroPerm[®] 30 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 30 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes. FluoroPerm[®] 30 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 30 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

CONTRAINDICATIONS (REASONS NOT TO USE)

FluoroPerm 30 rigid gas permeable contact lenses are contraindicated by the presence of any of the following conditions:

- Acute or subacute inflammations of the anterior segment of the eye.
- Any eye disease, injury, or abnormality, other than keratoconus, that affects the cornea, conjunctive or eyelids.
- Insufficiency of lacrimal secretion (dry eyes).

- Corneal hypoesthesia (reduced corneal sensitivity).
- Any systemic disease, which may affect the eye or be exacerbated by wearing contact lenses.
- Allergic reactions of ocular surfaces or adnexa which may be induced or exaggerated by wearing contact lenses and/or using contact lens solutions.
- Any active corneal infection (bacterial, fungal or viral).

WEARING SCHEDULE

THE EYE CARE PRACTITIONER SHOULD DETERMINE THE WEARING SCHEDULE. Patients tend to over wear the lenses initially. It is important to adhere to the initial maximum wearing schedule. Regular checkups, as determined by the eye care practitioner, are also extremely important.

The maximum suggested wearing time for FluoroPerm 30 rigid gas permeable contact lenses is:

DAILY WEAR (less than 24 hours while awake)

DAY	1	2	3	4	5	6	7	8	9	10 - 14	15 & after
SUGGESTED HOURS	3	4	5	6	7	8	9	10	11	12	All waking hours
HOURS WORN											

DO NOT SLEEP WHILE WEARING YOUR FLUOROPERM 30 RIGID GAS PERMEABLE CONTACT LENS. Studies have not been completed to show that the FluoroPerm 30 rigid gas permeable contact lens is safe to wear during sleep. There is a tendency for some patients to overwear the lenses initially. It is important to adhere to the maximum wearing schedule above. Regular check-ups, as determined by the eye care practitioner, are extremely important.

FLUOROPERM® 151
(paflucocon D)

RIGID GAS PERMEABLE
CONTACT LENSES FOR DAILY WEAR AND EXTENDED WEAR

Spherical contact lenses for:
Nearsightedness (myopia).
Farsightedness (hyperopia).

DESCRIPTION

FluoroPerm® 151 (paflucocon D) rigid gas permeable contact lenses for daily wear and extended wear are available as lathe cut or molded firm contact lenses with spherical front and back surfaces in tinted versions. The posterior curve is selected so as to properly fit an individual eye and the anterior curve is selected to provide the necessary optical power to correct refractive error. A peripheral curve system on the posterior surface allows tear exchange between the lens and the cornea.

The FluoroPerm 151 rigid gas permeable contact lens material is a thermoset copolymer derived from fluorosilicone acrylate monomers.

FluoroPerm 151 rigid gas permeable tinted lenses offer a handling aid for locating the lens. These products may be plasma treated.

The lenses have the following attributes.

FLUOROPERM[®] 151 (paflucocon D)

Refractive Index	1.442(Nd at 25°C)
Luminous Transmittance ⁺ (Blue)	93%
Luminous Transmittance (Crystal Blue)	98%
Wetting Angle (Receding Angle) ⁺⁺	42°
Wetting Angle (Receding Angle) ⁺⁺⁺	70°
Specific Gravity	1.10
Hardness (Shore D)	79
Water Content	<1%
Oxygen Permeability*	151 x 10 ⁻¹¹ Dk at 35°C
Oxygen Permeability**	100 x 10 ⁻¹¹ Dk at 35°C

⁺ Determination of the Spectral and Luminous Transmittance, ISO 8599:1994

⁺⁺ Adapted from: A New Method for Wetting Angle Measurement; Madigan, et. al., International Eyecare, 01/1998, vol. 2, no. 1, p. 45

⁺⁺⁺ Sessile Drop Technique per ANSI Z80.20. 8.11

* (cm²/sec) (mL O₂) / (mL x mm Hg) Revised Method of I. Fatt

** (cm²/sec) (mL O₂) / (mL x mm Hg) ISO/ANSI Fatt Method

Lens Parameters:

Chord Diameter	7.0 to 11.5 mm
Center Thickness	0.05 to 0.70 mm
Base Curve	6.50 to 9.00 mm
Powers Daily Wear	-20.00 to +12.00 Diopters
Powers Extended Wear	-20.00 to +8.00 Diopters

ACTION

FluoroPerm 151 rigid gas permeable contact lenses are intended for daily wear or extended wear. When placed on the human cornea, the FluoroPerm 151 rigid gas permeable contact lens acts as a refracting medium to focus light rays upon the retina.

INDICATIONS (USES)

FluoroPerm[®] 151 rigid gas permeable spherical or aspheric contact lenses are indicated for daily wear and extended wear from 1 to 7 days between removals for cleaning and disinfection as recommended by the eye care practitioner. FluoroPerm[®] 151 rigid gas permeable bifocal, or toric contact lenses are indicated for daily wear only.

FluoroPerm[®] 151 rigid gas permeable spherical, aspheric and bifocal contact lenses are indicated for the correction of visual acuity in not-aphakic persons with nondiseased eyes that are nearsighted (myopic), farsighted (hyperopic), and may exhibit corneal astigmatism up to 4.00 diopters that does not interfere with visual acuity. FluoroPerm[®] 151 contact lenses are indicated for persons requiring keratoconus management with otherwise non-diseased eyes, daily wear application only. FluoroPerm[®] 151 toric contact lenses are indicated to correct astigmatism of up to 6.00 diopters. FluoroPerm[®] 151 bifocal lenses are indicated to treat presbyopia up to +4.00 D add power.

CONTRAINDICATIONS (REASONS NOT TO USE)

FluoroPerm 151 rigid gas permeable contact lenses are contraindicated by the presence of any of the following conditions:

- Acute or subacute inflammations of the anterior segment of the eye.

- Any eye disease, injury, or abnormality, other than keratoconus, that affects the cornea, conjunctive or eyelids.
- Insufficiency of lacrimal secretion (dry eyes).
- Corneal hypoesthesia (reduced corneal sensitivity).
- Any systemic disease, which may affect the eye or be exacerbated by wearing contact lenses.
- Allergic reactions of ocular surfaces or adnexa which may be induced or exaggerated by wearing contact lenses and/or using contact lens solutions.
- Any active corneal infection (bacterial, fungal or viral).

WEARING SCHEDULE

THE EYE CARE PRACTITIONER SHOULD DETERMINE THE WEARING SCHEDULE. Patients tend to over wear the lenses initially. It is important to adhere to the initial maximum wearing schedule. Regular checkups, as determined by the eye care practitioner, are also extremely important.

The maximum suggested wearing time for FluoroPerm 151 rigid gas permeable contact lenses is:

DAILY WEAR (less than 24 hours while awake)

DAY	1	2	3	4	5	6	7	8	9	10 - 14	15 & after
SUGGESTED HOURS	3	4	5	6	7	8	9	10	11	12	All waking hours
HOURS WORN											

EXTENDED WEAR

Not every patient is able to wear the FluoroPerm 151 rigid gas permeable contact lens on an extended wear basis, even if able to wear the lens for daily wear. An initial 14-day daily wear period is recommended prior to overnight wear. Regular check-ups, as determined by the eye care practitioner, are extremely important. The lens should be removed at least once every week (7 days) for cleaning and disinfection.

Extended wear lenses should not be used to correct keratoconus.

DAY	SUGGESTED HOURS	HOURS WORN
15 (and after)	24 hours	

WARNING

The risk of ulcerative keratitis has been shown to be greater among wearers of extended wear lenses than among wearers of daily wear lenses. The risk among extended wear lens wearers increases with the number of consecutive days that lenses are worn between removals, beginning with the first overnight use. This risk can be reduced by carefully following directions for routine lens care, including cleaning of the lens storage case. Additionally, smoking increases the risk of ulcerative keratitis for contact lens wearers.

It is recommended that contact lens wearers see their eye care practitioners twice each year or, if directed, more frequently.

GENERAL INFORMATION

Convention: Reference to FluoroPerm rigid gas permeable contact lenses indicates all four materials – FluoroPerm[®] 92 (paflucocon A), FluoroPerm[®] 60 (paflucocon B), FluoroPerm[®] 30 (paflucocon C), and FluoroPerm[®] 151 (paflucocon D).

TINTS

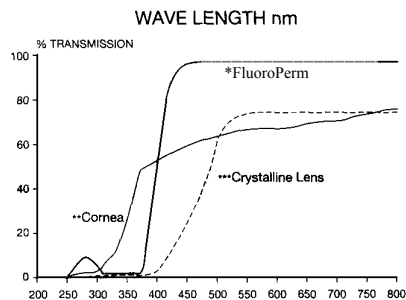
FluoroPerm rigid gas permeable contact lenses are available in untinted (clear) and tinted [blue, crystal blue, majestic blue, brown (daily wear only), gray, and green] versions with or without ultraviolet absorber. The tinted lenses contain one or more of the following color additives: D&D Green No. 6, D&C Red No. 17, D&C Violet No. 2 and Perox Yellow No. 9. NOTE: Not all materials are available in all colors or with ultraviolet absorber.

UV ABSORBER

FluoroPerm rigid gas permeable contact lenses are available with an ultraviolet absorber. The ultraviolet absorber, Uvinul D-49, has been integrated as an additive within the FluoroPerm polymer matrix and blocks up to 97% of light below 380 nm. The UV absorber is 2,2'-dihydroxy-4,4'-dimethoxybenzophenone.

WARNING: UV-absorbing contact lenses are not substitutes for protective UV-absorbing eyewear such as UV-absorbing goggles or sunglasses. Persons should continue to use their protective UV-absorbing eyewear as directed.

NOTE: The effectiveness of wearing UV-absorbing contact lenses in preventing or reducing the incidence of ocular disorders associated with exposure to UV-light has not been established at this time.



*FluoroPerm – 0.10 mm thick; blue

**CORNEA - Human cornea from a 24-year old person as described in Lerman, S., Radiant Energy and the Eye, MacMillan, New York, 1980, p. 58, figure 2-21.

***CRYSTALLINE LENS - Human crystalline lens from a 25-year old person as described in Waxler, M., Hitchins, V.M., Optical Radiation and Visual Health, CRC Press, Boca Raton, Florida, 1986, p.19, figure 5.

WARNINGS

PROBLEMS WITH CONTACT LENSES AND LENS CARE PRODUCTS COULD RESULT IN SERIOUS INJURY TO THE EYE. It is essential that you follow your eye care practitioner's directions and all labeling instructions for proper use of your contact lenses and lens care products. **EYE PROBLEMS, INCLUDING CORNEAL ULCERS, CAN DEVELOP RAPIDLY AND LEAD TO LOSS OF VISION; THEREFORE, IF YOU EXPERIENCE EYE DISCOMFORT, EXCESSIVE TEARING, VISION CHANGES, OR REDNESS OF THE EYE, OR OTHER PROBLEMS WITH YOUR EYES, IMMEDIATELY REMOVE YOUR LENSES, AND PROMPTLY CONTACT YOUR EYE CARE PRACTITIONER.**

PRECAUTIONS - PRACTITIONER

Clinical studies have demonstrated that contact lenses manufactured from the FluoroPerm rigid gas permeable contact lens material are safe and effective for their intended use. However, the clinical studies may not have included all design configurations or lens parameters that are presently available in the materials. Extended wear lenses should not be used to correct keratoconus.

Consequently, when selecting an appropriate lens design and parameter, the eye care practitioner must consider all factors that affect lens performance and ocular health. The potential impact of these factors must be weighed against the patient's needs, therefore, the continuing ocular health of the patient and lens performance on the eye should be carefully monitored.

Each FluoroPerm lens is supplied nonsterile in an individual plastic case. The lens is shipped dry; or, wet shipped in Unique-pH[®] Multi-Purpose Solution. This solution contains hydroxypropyl guar, a unique wetting and conditioning polymer system, polyethylene glycol, Tertronic*, boric acid, propylene glycol; and, is preserved with POLYQUAD[®] (polyquarternium-1) 0.0011% and edetate disodium 0.01%. If the patient has experienced a prior history of allergy to any of these ingredients, remove the lens from the solution and soak the lens 24 hours in unpreserved saline prior to cleaning, disinfecting and dispensing.

Unique-pH[®] is a Trademark of Alcon Laboratories, Inc.

* Registered Trademark of BASF Corp.

Never reuse the solution. You may store the lenses in the unopened container until ready to dispense, up to a maximum of thirty (30) days from the Fill Date (see container). When a lens has been stored for 30 days in its original packaging solution, it should be cleaned and disinfected with an approved product (see recommended product list in the Lens Care Directions section), and placed into inventory as you presently do with any other RGP lens held in your office. Follow the directions on the selected disinfecting solution regarding prolonged storage.

PRECAUTIONS - PATIENT

Follow the instructions below to prevent damage to your eyes or to your lenses.

- Before you leave your practitioners office be able to promptly remove your lens or have someone else be able to remove your lens for you.
- **DO NOT WEAR YOUR FLUOROPERM RIGID GAS PERMEABLE CONTACT LENSES WHILE SLEEPING UNLESS YOUR WEARING SCHEDULE IS INTENDED FOR EXTENDED WEAR.**
- Always wash your hands with an additive free soap, rinse thoroughly and dry on a lint free towel before you handle your lenses. Eye irritation may result if cosmetics, lotions, soaps, creams and deodorants come in contact with your lenses and if the lenses are contaminated by infectious or non-infectious debris.
- Always follow the recommended lens care system for your FluoroPerm lenses. Use the recommended lens care solutions and carefully follow the recommended directions.
- Always use FRESH rinsing, disinfecting and storage solutions.
- Do not use saliva or anything other than the recommended solutions to wet your lenses.
- Always keep the lenses completely immersed in the recommended storage solution when the lenses are not being worn.
- Avoid using aerosol products such as hair spray while wearing your lenses. If hair sprays are used, keep your eyes closed until the spray has settled, otherwise, the lenses may be damaged.
- Avoid all harmful or irritating vapors and fumes while wearing your lenses.

- Do not swim with your lenses in place.
- Never use tweezers or other tools to remove your lens from the lens container. Do not touch the lens with your fingernails.
- Always inform your doctor (general health care practitioner) that you wear contact lenses.
- Always consult your eye care practitioner before using any medicine in your eyes.
- Always inform your employer that you wear contact lenses. Some jobs may require the use of protective eye equipment or may require that you not wear contact lenses.
- As with any contact lens, follow-up visits are necessary to assure continued health. **CHECK WITH YOUR EYE CARE PRACTITIONER.**
- To minimize lens warpage during cleaning, the lenses should be cleaned in the palm of the hand rather than between the thumb and fingers.
- Do not heat the conditioning solution and lenses.
- The safety of these lenses with medications or contact lens solutions other than those recommended has not been established.
- If your lens sticks (stops moving) on the eye, follow the recommended directions for "Care for a Sticking Lens". The lens must move freely on the eye for the continued health of the eye. If nonmovement of the lens continues, immediately consult your eye care practitioner.
- **CAUTION:** Nonsterile. Clean and condition lenses prior to use.

ADVERSE EFFECTS (PROBLEMS AND WHAT TO DO)

The following problems may occur.

- Eyes sting, burn or itch (irritation)
- Comfort is less than when lens was first placed on eye
- Feeling of something in the eye (foreign body, scratched area, abrasion)
- Excessive pain
- 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 you notice any of these adverse effects, **IMMEDIATELY REMOVE YOUR 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 your eye. Place the lens in the storage case and contact your eye care practitioner.
- If the lens has dirt, an eyelash, or other foreign body on it, or the problem stops and the lens appears undamaged, thoroughly clean, rinse and disinfect the lenses; then reinsert it.
- If the problem continues, **IMMEDIATELY** remove your contact lens and consult your eye care practitioner.

When any of the above symptoms occur, a serious condition such as infection, corneal ulcer, neovascularization, iritis, persistent stromal edema or GPC (giant papillary conjunctivitis) may be present.

Immediately remove your lenses and seek professional identification of the problem and prompt treatment to avoid serious eye damage, including corneal scarring, opacification, blindness or loss of eye.

FITTING

Conventional methods of fitting rigid contact lenses apply to the FluoroPerm rigid gas permeable contact lens. For a description of fitting techniques, refer to the Fitting Guide for FluoroPerm Rigid Gas Permeable Contact Lenses. Copies of which are available from:

Paragon Vision Sciences
947 E. Impala Avenue
Mesa, Arizona 85204-6619

CAUTION: Federal (US) law restricts this device to sale by, or on the order of a licensed practitioner.

LENS CARE DIRECTIONS

Always wash your hands with an additive-free soap, rinse thoroughly and dry on a lint-free towel before you handle your contact lenses.

Never use tweezers or other tools to remove your lens from the lens container. Pour the lens into your hand.

FluoroPerm rigid gas permeable contact lenses must be both cleaned and disinfected each time you remove them. One procedure does not replace the other. Cleaning is necessary to remove mucus and film from the lens surface. Disinfecting is necessary to destroy harmful germs. Leave the FluoroPerm rigid gas permeable contact lenses in a storage solution for a minimum of 4 hours or as indicated on the product label. To minimize lens warpage during cleaning, the lenses should be cleaned in the palm of the hand rather than between the thumb and fingers.

Clean one lens first. The recommended procedure is to always clean the same lens first to avoid mix-ups. Rinse the lens thoroughly to remove the cleaning solution. Place the lens into the correct storage chamber and fill the chamber with the recommended disinfection system as recommended by your eye care practitioner. Clean and rinse the other lens in the same manner and place it in its chamber.

Tightly close the top of each chamber of the lens storage case.

To disinfect your lenses, leave them in the solution for at least 4 hours, or as indicated on the product label.

Do not heat the conditioning solution and lenses.

Leave the lenses in the unopened storage case until you are ready to put them on your eyes.

After you remove your lens from the lens case, empty and rinse your lens storage case with fresh, hot running tap water and allow it to air dry. When you next use the case, refill it with fresh storage solution.

FluoroPerm rigid gas permeable contact lenses should be disinfected using only a chemical (not heat) disinfection system.

The following is a list of products available for use with FluoroPerm rigid gas permeable contact lenses. This is not an exclusive list. You may use other lens care solutions as recommended by your eye care practitioner.

SYSTEM PROCESS	CHEMICAL (not heat) DISINFECTION SYSTEM
Cleaning	Unique-pH [®] Multi-Purpose Solution, SupraClens [®] , Opti-Clean [®] II, ProFree/GP [®]
Disinfection	Unique-pH [®] Multi-Purpose Solution, Wet-N-Soak [®] Plus, Barnes-Hind [®] ComfortCare GP Wetting and Soaking Solution
Lubrication	Clerz [®] Plus

PRODUCT LIST

Unique-pH[®] Multi-Purpose Solution, SupraClens[®], Opti-Clean[®] II, Clerz[®] Plus by Alcon Laboratories, Inc.

ProFree/GP[®], Wet-N-Soak[®] Plus, Barnes-Hind[®] ComfortCare GP Wetting and Soaking Solution by Advanced Medical Optics (AMO)

Follow the instructions provided with each lens care solution. Failure to adhere to these procedures may result in the development of serious ocular complications. A patient should not switch from one care system to another unless the eye care practitioner has determined that this is necessary. Do not mix or alternate the disinfection and storage systems unless so indicated on the product label.

CARE FOR A STICKING LENS

If the lens sticks (stops moving) on the eye, apply a few drops of a lubricating solution. Wait until the lens begins to move freely on your eye before removing it. If nonmovement of the lens continues, immediately consult your eye care practitioner.

HOW SUPPLIED

Each FluoroPerm lens is supplied nonsterile in an individual plastic case. The lens is shipped dry; or, wet shipped in Unique-pH[®] Multi-Purpose Solution. This solution contains hydroxypropyl guar, a unique wetting and conditioning polymer system, polyethylene glycol, Tertronic*, boric acid, propylene glycol; and, is preserved with POLYQUAD[®] (polyquaternium-1) 0.0011% and edetate disodium 0.01%. The case, packing slip or invoice is marked with the base curve, dioptric power, diameter, center thickness, inclusion of UV absorber, lot number, fill date and the color of the lens.

Unique-pH[®] is a Trademark of Alcon Laboratories, Inc.

* Registered Trademark of BASF Corp.

Never reuse the solution. You may store the lens in the unopened container until ready to dispense, up to a maximum of thirty (30) days from the Fill Date. When a lens has been stored for 30 days in its original packaging solution, it should be cleaned and disinfected with an approved product (see recommended product list in the Lens Care Directions section), and placed into inventory as you presently do with any other RGP lens held in your office. Follow the directions on the selected disinfecting solution regarding prolonged storage.

For information on material specifications contact:

Paragon Vision Sciences
947 East Impala Avenue
Mesa, Arizona 85204-6619

1-800-528-8279
1-480-892-7602
1-480-926-7369 FAX

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