

October 20, 2020

CooperVision, Inc.
Marie Dutton
Senior Regulatory Affairs Specialist
6101 Bollinger Canyon Road, Suite 500
San Ramon, CA 94583

Re: K202756

Trade/Device Name: Clariti 1 day (somofilcon A) Soft (Hydrophilic) Daily Disposable Contact Lens

with UV blocker

Regulation Number: 21 CFR 886.5925

Regulation Name: Soft (Hydrophilic) Contact Lens

Regulatory Class: Class II Product Code: LPL, MVN Dated: September 18, 2020 Received: September 21, 2020

Dear Marie Dutton:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for

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devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to https://www.fda.gov/medical-device-problems.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

J. Angelo Green, Ph.D.
Assistant Director
DHT1A: Division of Ophthalmic Devices
OHT1: Office of Ophthalmic, Anesthesia,
Respiratory, ENT and Dental Devices
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120

Expiration Date: 06/30/2023 See PRA Statement below.

510(k) Number (if known)	
K202756	
Device Name	
Clariti 1 day (somofilcon A) Soft (Hydrophilic) Daily Disposable Contact Lens with UV Blocker	

Indications for Use (Describe)

The CLARITI 1 DAY (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker is indicated for daily wear single use only for the correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 2.00 Diopters that does not interfere with visual acuity.

The CLARITI 1 DAY TORIC (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters.

The CLARITI 1 DAY MULTIFOCAL (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may require a reading addition of +3.00 Diopters or less and may exhibit astigmatism up to 1.50 Diopters or less.

The CLARITI 1 DAY MULTIFOCAL TORIC (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV blocker is indicated for daily wear single use only for the optical correction of refractive ametropia (myopia and hyperopia) and/or presbyopia in phakic or aphakic persons with non-diseased eyes that may exhibit astigmatism up to 10.00 Diopters and require a reading addition of +3.00 Diopters or less.

Type of Use (Select one or both, as applicable)	
Prescription Use (Part 21 CFR 801 Subpart D)	Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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510(k) Summary

I. SUBMITTER:

CooperVision, Inc. 6101 Bollinger Canyon Road, Suite 500 San Ramon, CA 94583

Contact Person:

Marie Dutton Senior Regulatory Affairs Specialist 6101 Bollinger Canyon Road, Suite 500 San Ramon, CA 94583

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Date Prepared:

September 18, 2020

II. DEVICE:

Trade Name: Clariti 1 day (somofilcon A) Soft (Hydrophilic)

Daily Disposable Contact Lens with UV Blocker

Common Name: Soft (hydrophilic) Contact Lens

Classification Name: Lens, Contact, (Disposable) [21 CFR 886.5925 (b) (1)]

Regulatory Class: II

Product Code: LPL, MVN Classification Panel: Ophthalmic

III. PREDICATE DEVICE:

CooperVision's Clariti 1 day (somofilcon A) Soft (Hydrophilic) Daily Disposable Contact Lens with UV Blocker, K130331

IV. DEVICE DESCRIPTION:

The device description for the subject device and predicate device is identical. The Clariti 1 day (somofilcon A) Soft (Hydrophilic) Daily Disposable Contact Lens with UV Blocker is available as sphere lens, toric lens, multifocal lens, and multifocal toric lens.

In its hydrated state, Clariti 1 day (somofilcon A) Soft (hydrophilic) Daily Disposable Contact Lens with UV Blocker when placed on the cornea acts as a refracting media to focus light rays on the retina.

Clariti 1 day (somofilcon A) Soft (hydrophilic) Contact Lenses for Daily Wear Single Use are a hydrophilic co-polymer of silicone containing monomers and hydrophilic monomers which is cross-linked with tetraethyleneglycol dimethacrylate and di-functional methacryloxypropyl-terminated poly(dimethylsiloxane).

When hydrated the lens consists of 44.0% somofilcon A and 56.0% water by weight of saline immersed in normal saline. A benzophenone UV absorbing monomer is used in the contact lens to help protect against transmission of harmful UV radiation and Clariti 1 day (somofilcon A) Soft contact lenses help protect against transmission of harmful UV radiation to the cornea and into the eye.

The average transmittance characteristics are less than 5% in the UVB range of 280 to 315nm and less than 50% in the UVA range of 316-380nm.

The lens has a hemispherical flexible shell, which covers the cornea and a portion of the adjacent sclera, with the following dimensions:

Chord Diameter: 13.0mm to 15.5mm
 Centre Thickness: 0.03mm to 0.50mm
 Base Curve: 7.5mm to 9.30mm
 Powers: -20.00 DS to +20.00 DS
 Toric Cylinder options: -0.75, -1.25, -1.75 and -2.25
 Toric Axis options: 10° to 180° (10° steps)

Multifocal Add:

Lens "LOW" = "low" for spectacle near ADD lens (Max +2.25 ADD) Lens "HIGH" = "high" for spectacle near ADD lens (+2.50 ADD or greater)

The physical/optical properties of the lenses are:

Refractive Index: 1.4003
 %Transmittance @ 590nm: 98.13
 %Transmittance @ 280-315nm: 0.71
 %Transmittance @ 316-380nm: 20.62
 Surface Character: Hydrophilic

• Water Content: 56%

• Oxygen Permeability (DK): 60 x 10⁻¹¹ (cm²/sec)

(ml O₂/ml x mmHg) at 35°C (Fatt Method for determination of oxygen

permeability)

• Specific Gravity: 1.17

V. INDICATIONS FOR USE:

The indications for use statement for the subject device and predicate device is identical.

Lens Design	Indication				
Sphere	The CLARITI 1 DAY (somofilcon A) Soft (hydrophilic) Daily				
	Disposable Contact Lens with UV blocker is indicated for daily wear				
	single use only for the correction of refractive ametropia (myopia and				
	hyperopia) in phakic or aphakic persons with non-diseased eyes that				
	may exhibit astigmatism up to 2.00 Diopters that does not interfere				
	with visual acuity.				
Toric	The CLARITI 1 DAY TORIC (somofilcon A) Soft (hydrophilic)				
	Daily Disposable Contact Lens with UV blocker is indicated for daily				
	wear single use only for the optical correction of refractive ametropia				
	(myopia and hyperopia) in phakic or aphakic persons with non-				
	diseased eyes that may exhibit astigmatism up to 10.00 Diopters.				
Multifocal	The CLARITI 1 DAY MULTIFOCAL (somofilcon A) Soft				
	(hydrophilic) Daily Disposable Contact Lens with UV blocker is				
	indicated for daily wear single use only for the optical correction of				
	refractive ametropia (myopia and hyperopia) and/or presbyopia in				
	phakic or aphakic persons with non-diseased eyes that may require a				
	reading addition of +3.00 Diopters or less and may exhibit astigmatism				
	up to 1.50 Diopters or less.				
Multifocal	The CLARITI 1 DAY MULTIFOCAL TORIC (somofilcon A) Soft				
Toric	(hydrophilic) Daily Disposable Contact Lens with UV blocker is				
	indicated for daily wear single use only for the optical correction of				
	refractive ametropia (myopia and hyperopia) and/or presbyopia in				
	phakic or aphakic persons with non-diseased eyes that may exhibit				
	astigmatism up to 10.00 Diopters and require a reading addition of				
	+3.00 Diopters or less.				

VI. COMPARISON OF TECHNOLOGICAL CHARACTERISTICS WITH THE PREDICATE DEVICE:

The technological principle for both the subject and predicate device is identical.

Technology/Material Comparison				
	Predicate Device	Subject Device		
Product Name	Clariti 1 day (somofilcon A) Soft (Hydrophilic) Daily Disposable Contact Lens with UV Blocker	Same		
Material USAN Name	somofilcon A	Same		
510(k) Number	K130331	Current Submission		
FDA Category (Group)	Materials having a Dk greater than 40 Dk units (using mmHg) and having a Dk greater than that expected based on the materials' water content alone (Group V)	Same		
Manufacturing Method	Cast Molding	Same		
Wearing and Replacement Schedule	Daily Wear Single Use	Same		
Sterilization Method	Moist Heat	Same		
Packaging Materials	Injection molded polypropylene blisters covered by aluminum foil laminate and blister strips are packed into printed cartons	Same		
Packaging Solution	Borate Buffered Saline Solution containing 0.005% w/v Poloxamer 407	Same		
Blue Visibility Tint	No	Same		
Tint	None	Same		
UV Blocker	UV416	Same		

VII. PERFORMANCE DATA:

The performance specifications/parameters for both the subject and predicate device are identical.

Performance Specifications/Parameters Comparison				
	Predicate Device	Subject Device		
	Clariti 1 day	Clariti 1 day		
	(somofilcon A)	(somofilcon A)		
	K130331	Current Submission		
Water Content (%)	56	Same		
Refractive Index	1.401	Same		
Oxygen Permeability	$60 \times 10^{-11} [(cm^2/sec) \times (ml)]$	Same		
(Dk @ 35°C)	O_2 /(ml x mm Hg)]			
Base Curve (mm)	8.6	Same		
Diameter (mm)	14.0	Same		
Light Transmittance (%)	>95	Same		
Modulus (MPa)	≥0.3	Same		
Tensile Strength (MPa)	≥0.4	Same		
Elongation to Break (%)	≥100	Same		
Surface Treatment	No	Same		
Center Thickness (mm)	Varies with power	Same		
Power Range (D)	-20.00 to + 20.00	Same		

VIII. CONCLUSIONS:

This 510(k) is submitted in accordance with the May 12, 1994 Premarket Notification [510(k)] Guidance Document for Daily Wear Contact Lenses, amended June 28, 1994, for the change from a biological release to parametric release of finished product. The modification does not warrant performance testing to support substantial equivalence. Therefore, the subject device is considered substantially equivalent to the predicate device.