DECLARATION OF CONFORMITY						
(check all additional conformity route(s) based on EU MDD Article 11 requirements for the device class and specifics)						
Annex II (4) □ Ann	ex V □ Annex III □	Annex VII □				
Annex II (3) ⊠ Ann	ex VI □ Annex IV □					
Technical File Number and Version: DU-VC-004 Version 07						
Device Trade Name: DAILIES AquaComfort Plus (nelfilcon A) Soft Contact Lens						
Supersedes (Date): 28-Oct-2020						
Manufacturer: Alcon Laboratories, Inc.		ative in the European Community*:				
Address: 6201 South Freeway, Fort Worth, TX 76134-2099, USA	Alcon Laboratories Bel	gium , 2870 Puurs-Sint-Amands, Belgium				
Manufacturing Site(s):			, Deiglain			
	*Previously Alcon Labor	ratories (UK) Ltd. , Frimley, Camberley Surrey, GU16				
CIBA VISION GmbH Industriering 1,63868 Grosswallstadt, Germany	7SR, United Kingdom					
Alcon Research, LLC 11440 Johns Creek Parkway, Duluth, GA 30097, U	JSA Alcon Laboratories Bel Rijksweg 14, 2870 Puu					
CIBA VISION Asian Manufacturing and Logistics Pte Ltd. 133 Tuas South Avenue 3, Singapore 637550, Singapore						
Device (Trade Name)	GMDN Code & Term	Catalogue Number	Class			
DAILIES AquaComfort Plus						
DAILIES AquaComfort Plus Toric	47841 Soft Corrective Contact Lens,	N/A	lla			
DAILIES AquaComfort Plus Multifocal	Daily-disposable					
DAILIES AquaComfort Plus Asphere**						
The device(s) covered by this declaration are in conformity with the European Medical Devices Directive 93/42/EEC as well as other, relevant Union legislation that make provisions for the issuing of a declaration of conformity as listed.						
Alcon Laboratories, Incorporated hereby declares under its sole responsibility that the listed device(s) and Quality Systems						
conform(s) to: EU MDD 93/42/EEC as amended						
This Declaration is applicable to all products listed and released after the Date of Issuance of this Declaration of Conformity, and until a revised Declaration of Conformity is issued.						
Notified Body Information: Applicable ⊠ Not Applicable □						
Conformity Assessment Certificate Number(s): G1 020895 0393 Conformity Certificate Validity Period: 05-Feb-2021 to 26-May-2024						
Notified Body: TÜV SÜD Product Service GmbH*** ***PreviouslyBSI, Kitemark Court, Davy Avenue, Knowhill, Milton Kaynes MK5 8PP UK with identification number 0086.						
Identification number: 0123						
Address: Ridlerstraße 65, 80339 Munich, Germany						
Regulations, Directives and Standards Applied: EN ISO 13485 as currently published						



Place of Issue: Alcon Laboratories, Incorporated, Fort Worth, TX, USA	Date of Issue: 03-Mar-2021	Lakota, Sherri	Digitally signed by Lakota, Sherri DN: cn=Lakota, Sherri, ou=PDF Internal Signing, o=Alcon Vision, LLC Reason: I am approving this document Date: 2021.03.03 10:5429-06'00' Adobe Acrobat DC version: 2015.006.30503
		Name/Title/Function/Date: Sherri Lakota / VP GRA VC & DEOH	