DECLARATION OF CONFORMITY			
(check all additional conformity route(s) based on EU MDD Article 11 requirements for the device class and specifics)			
- ()	nex V □ Annex III □ nex VI □ Annex IV □	Annex VII □	
Technical File Number and Version: DU-VC-003 Version 04 Device Trade Name: AIR OPTIX Night & Day AQUA (lotrafilcon A) Soft Contact Lens Supersedes (Date): 10-Dec-2020			
Manufacturer: Alcon Laboratories, Inc. Address: 6201 South Freeway, Fort Worth, TX 76134-2099, USA	Authorized Representative in the European Community*: Alcon Laboratories Belgium Address: Lichterveld 3 2870 Puurs-Sint-Amands, Belgium		nunity*:
Manufacturing Site(s): CIBA VISION Johor Sdn. Bhd. No. 1 Jalan DPB/5, Pelabuhan Tanjung Pelepas, Gelang Patah, Johor Darul Takzim, Johor 81560, Malaysia	*Previously Alcon Labora		y, GU16
PT CIBA VISION Batam JL Beringin Lot #204 Batamindo Industrial Park, Muka Kuning, Batam Island 29433, Indonesia			
Device (Trade Name)	GMDN Code & Term	Catalogue Number	Class
AIR OPTIX Night & Day AQUA (lotrafilcon A)	47843 Soft Corrective Contact Lens, 36054 Therapeutic Contact Lens, Extended wear	N/A	llb
AIR OPTIX Night & Day AQUA (lotrafilcon A) The device(s) covered by this declaration are in cother, relevant Union legislation that make provis	36054 Therapeutic Contact Lens, Extended wear onformity with the European Medical Devices	Directive 93/42/EEC as w	
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Place of Issue:
Alcon Laboratories,
Incorporated, Fort
Worth, TX, USA

Date of Issue:

Lakota,
Digitally signed by Lakota, Sherri
DN: cn=Lakota, Sherri, ou=PDF Internal
Signing, o=Alcon Vision, LLC
Reason: I am approving this document
Date 2012.03.31 10:53:27 -0600'
Adobe Acrobat DC version:
2015.006.30503

Name/Title/Function/Date: Sherri Lakota / VP GRA VC & DEOH