Documents provided by us based on your request, present the sole property of Bausch Health/BAUSCH+LOMB, or the respective manufacturer and can be used solely for the designated purposes in accordance with by the Regulation (EU) 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and of the Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament and Council of 5 April 2017 11 5 the European Faciliament April 2017 11 5 the Euro

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Rochester, NY 14609 USA 665,330,6000

EU Declaration of Conformity

In accordance with Annex IV of Regulation (EU) 2017/745 of the European Parliament and of the Council

Technical Documentation Number	745-002
Legel Manufacturer	Bausch & Lomb Incorporated 1400 North Goodman St. Rochester, NY 14609 USA SRN: US-MF-000001029
European Authorized Representative	Bausch & Lomb Incorporated Cork Road Industrial Estate Waterford, X91 V983, Ireland SRN: IE-AR-000000094
Notified Body	Nalional Standards Authority of Ireland (NSAI) 1 Switt Square Northwood, Santry Dublin 9, Ireland Noblied Body Number: 0950
Products	Bausch + Lorid UKTRA® ONE DAY (kalillicon A) Copied Lenses
Product Code(s)	KAPBSDD
Baeic UDI-DI	8310119B0020YV
Global Medical Devices Nomenclatura Code	47841 - Soft corrective contact lons, daily-
Term Nama	Øjsposable
EMDN Code	QQ21004010101
Classification	Class IIa (Annex VIII, Rule 5)
Conformity Assessment Probedure	Regulation (EU) 2017/745 Annex IX Conformity assessment based on a quality management system and on assessment of technical documentation
Intended Purpose	Bausch + Lomb kall/licon A Conlect Lens is Indicated for the deily wear correction of retractive ametropia (myopia and hyperopia) in aphakic and/or non-ephakic persons with non-diseased syes that exhibit refractive astigmalism of 2.00 diopters or less, that does not interfere with visual acuity. The lens is to be prescribed for single-use disposable wear and is to be disparded effer each removal.

Documents provided by us based on your request, present the sole property of Bausch Health/BAUSCH+LOMB, or the respective manufacturer and can be used solely for the designated purposes in accordance with by the Regulation (EU) 2017 45 5 the Enrope and alliament and of the Council of 5 April 2017 100 100 100 degices as amended. Hochester, NY 14609 USA

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This EU Declaration of Conformity is issued under the sole responsibility of the Legal Manufacturer.

Wa, Bausch & Lomb Incorporated, hereby declare the above listed Medical Devices comply with Medical Device Regulation (EU) 2017/745. Above products are developed and manufactured in compliance with the EU MDR and the applicable European harmonized standards and Common Specifications as applicable.

This declaration is made on the basis of EU Quality System Cartificate Number MD19.1854 and EU MDR EC Certificate Number 745.002, issued by the Notified Body stated above, in accordance with Annex IX, Chapters I and III of Medical Device Regulation (EU) 2017/745 and the products above comply with Annex L.

Place of Issue; Refer to Legal Menulecturer's Address above.

Signature:

Date:

Name/Title/Position:

585 338,8000