Johnson Johnson VISION

CERTIFICATION OF COPY

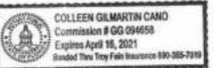
On this 29th day of March 2019, I certify that the attached is a true, exact and unaltered photocopy provided to me of:

Declaration of Conformity – senofilcon C – Version 5.0 – 2 pages

presented to me by Rosalind D. Williams, Project Leader, Regulatory Affairs, and to the best of my knowledge, that the photocopied documents are neither a vital record nor a public record, certified copies of which are available from an official source other than a Notary Public.

United States of America	1
State of Florida	1
County of Duval	1

(Notary Public





VISION CARE, INC.

EU Technical File Version 5.0

DECLARATION OF CONFORMITY

Manufacturer	Johnson & Johnson Vision Care, Inc. 7500 Centurion Parkway Jacksonville, Florida 32256 United States
Product Name	senofilcon C Contact Lenses
Description	Spherical contact lenses are intended for Daily Wear for the optical correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may have 1.00D or less of astigmatism. Toric contact lenses are intended for Daily Wear for the optical correction of refractive ametropia (myopia and hyperopia) in phakic or aphakic persons with non-diseased eyes that may have astigmatism. These lenses have UV Blocking to help provide protection against transmission of harmful UV radiation to the comea and into the
Product Identification	eye. See page 2 of this Declaration
Classification	Ha
Classification Rationale	
	Rule 5
Declaration	This declaration of conformity is issued under the sole responsibility of the manufacturer, per Decision No 768/2008/EC. We, being the manufacturer/distributor within the European Economic Area, declare that the products covered by this declaration, documented in Technical File – senofilcon C Version 5.0, dated 26 March 2019, conform with the essential requirements and provisions of European Council Directive 93/42/EEC.
	We, the manufacturer/distributor, have been subject to the conformity procedures laid down in Annex II under the supervision of the British Standards Institution, a Notified Body authorized by the Netherlands Competent Authority, carrying the Notified Body Number 2797 and no application has been lodged with any other Notified Body.
	This declaration is supported by the Johnson & Johnson Vision Care, Inc. Quality Management Systems approved by EC Certificate for Quality Assurance Certificate Number CE 00387.

senofilcon C EU Technical File Version 5.0 – Declaration of Conformity Johnson & Johnson Vision Care, Inc. (JJVCI)

the following sites; Johnson & Johnson Vision Care, Inc. 7500 Centurion Parkway Jacksonville, Florida 32256 United States
Johnson & Johnson Vision Care European Vision Centre 8 Hanworth Road Sunbury TW16 5LN United Kingdom
AMO Ireland Block B, Liffey Valley Office Campus, Quarryvale, Co. Dublin D22 X0Y3 Ireland
The following are included: Product Family: ACUVUE® VITA™ Models: ACUVUE® VITA™ Brand Contact Lenses ACUVUE® VITA™ Brand Contact Lenses for ASTIGMATISM Product Listing includes Diagnostic, Revenue and Kit Configurations. GMDN Codes: 47842, Soft corrective contact lens, daily wear

Catherine Dillon

Director, Regulatory Affairs Johnson & Johnson Vision Care, Inc. Jacksonville, Florida 32256, USA

Thonas Wilbinson

Thomas Wilkinson Director, Quality Systems Johnson & Johnson Vision Care, Inc. Jacksonville, Florida 32256, USA 26 March 2019 26 March 2019

Date

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