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Document No. DOCAL-ScanEU12

DECLARATION OF CONFORMI

Manufacturer's name	NIDEK Co., Ltd.	SRN	Not issued yet	
Manufacturer's address	34-14 Maehama, Hiroishi	-cho, Gamago	ri, Aichi 443-0038, Ja	pan
European Representative name	NIDEK S.A.	SRN	FR-AR-0000003	341
European Representative address	Ecoparc, rue Benjamin Fra	ınklin, 94370 S	ucy En Brie, FRANCE	3
Identification of device	OPTICAL BIOMETER			
Model No.	AL-Scan			
Classification(Annex VIII, MDR)	IIa (Rule 10)			
Category (for RoHS)	8			
Basic UDI-DI	4987669210G3			

We herewith declare under our sole responsibility that the above mentioned products meet the provisions of the following EC Council Directive(s)/Regulation. All supporting documentation is retained under the premises of the manufacturer. The supporting technical documentation is also retained at NIDEK S.A., Ecoparc, rue Benjamin Franklin, 94370 Sucy En Brie, FRANCE.

General applicable directive(s)/regulation	Notified Body	Date CE Marking was affixed
REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT	DEKRA Certification B.V. Meander 1051, 6825 MJ Arnhem	May 11, 2012
AND OF THE COUNCIL of 5 April 2017 on medical devices	P.O. Box 5185, 6802 ED Arnhem, The Netherlands	CE
	Certificate No.: 4202074CE01 (Annex IX, Section 2 of MDR)	0344
COUNCIL DIRECTIVE 2011/65/EU	N/A	March 27, 2014
concerning restriction of the use of certain hazardous substances.		CE

Place: Aichi, Japan

Effective date: November 21, 2022

Signed by

Katsuaki Tohyama Senior Manager

Person responsible regulatory compliance

for

NIDEK Co., Ltd.

Date of signature: November 16, 2022