

# IDCP

## EC DECLARATION OF CONFORMITY

We,

IDCP BV  
Manuscriptstraat 12-14  
1321 NN Almere  
The Netherlands  
SRN: NL-MF-000000755

hereby declare under our sole responsibility that the CE-marked products to which this declaration relates,

CapillaryScope 500 (type number MEDL4N5, Basic UDI-DI  
87202991629MEDL4N5SS)  
CapillaryScope Pro 200 (type number MEDL4N Pro, Basic UDI-DI  
87202991629MEDL4NPro4T)  
and  
CapillaryScope Pro 500 (type number MEDL4N5 Pro, Basic UDI-DI  
87202991629MEDL4N5ProSJ)

having the intended purpose: The CapillaryScope is intended to make images of microcapillary vessels,

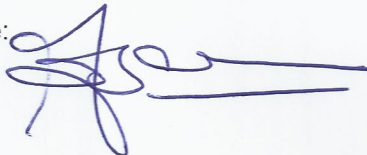
and have been classified as Class I, according to Annex VIII, Rule number 10, and the related DinoCapture software is classified as Class I, because the software drives the CapillaryScope or influences the use of CapillaryScope and consequently shall fall within the same class as the CapillaryScope,

and are in conformity with the General Safety and Performance Requirements of Regulation (EU) 2017/745 of the European Parliament and of the Council of 5 April 2017 on medical devices,

and are in conformity with the standards EN 1041:2008 and EN ISO 15223-1:2016,

and are in conformity with the requirements of directive 2011/65/EU of the European Parliament and of the Council of 8 June 2011 on the restriction of the use of certain hazardous substances in electrical and electronic equipment.

Signature:



Naarden, The Netherlands

Date: 9-2-2021

Name: Jan Boers

Function: Managing Director

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