## EU Declaration of Conformity

In accordance with the Regulation (EU) 2017/745 on medical devices (MDR 2017/745)

Manufacturer's Name: ILLUCO Corporation Ltd.

Manufacturer's Address: 102-304 SK Ventium, #166 Gosan-ro

Gunpo-si, Gyeonggi-do 15850

Republic of Korea

**SRN** (Single Registration

Number):

KR-MF-000012082

**Authorized Representative** 

Name (if applicable):

DermoScan GmbH

**Authorized Representative** 

Address (if applicable):

Ohmstraße. 1

93055 Regensburg, Germany

**Basic UDI-DI:** 88001906IDS1100BK

**Description of the Device(s):** Dermatoscope IDS-Series

Class I, non-sterile and non-measuring medical devices

Name of Device(s): IDS-1100, IDS-1100C, IDS-1000, IDS1000 PLUS, IDS-3100

**Risk Class:** Medical Device Class I

Not required for medical device class I **Notified Body:** 

**Conformity assessment route:** The procedures according to the Regulation (EU) 2017/745 on

medical devices:

Class I: EU conformity declaration according to Annex I,

Annex II, Annex III, and Annex XIV.

References to the relevant harmonized standards or

specifications in relation to this

conformity:

ISO 13485 Medical devices – Quality management system –

Requirements for regulatory purposes

EN 60601-1-2 Medical electrical equipment – Part 1-2 General

requirements for basic safety and essential performance -Collateral Standard: Electromagnetic disturbances -

Requirements and tests

This declaration of conformity is issued under the sole responsibility of ILLUCO Corporation Ltd. We hereby declare that the medical device(s) specified above meet the provision of the **Regulation** (EU) 2017/745 on medical devices (MDR 2017/745).

Signature. LTD

Place and date (dd.mm.yyyy) of issue:

PRESIDENT / ILPYO, HONG

Gunpo, Republic of Korea, 25.08.2021

Name: Ilpyo Hong Function: CEO