DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 93/42/EEC OF 14 JUNE 1993 CONCERNING MEDICAL DEVICES

MANUFACTURER: Shenzhen IMDK Medical Technology Co., Itd C Zone, 10F, Building 16, Yuanshan Industrial B Area, Gongming Street, Guangming District, 518106, Shenzhen, China.

MEDICAL DEVICE: PULSE OXIMETER, C101H1/C101A2/C101A3/C101B1/C101B2

CLASSIFICATION - ANNEX IX: CLASS IIA, RULE11

CONFORMITY ASSESSMENT ROUTE: ANNEX VII + V.3

We, <u>THE MANUFACTURER</u>, EXCLUSIVELY RESPONSIBLE FOR THE DECLARATION OF CONFORMITY, AND HEREWITH DECLARE THAT THE STATED MEDICAL DEVICES

MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE

93/42/EEC OF 14 JUNE 1993 CONCERNING MEDICAL DEVICES;
INCLUDING, AT 21 MARCH 2010, THE AMENDMENTS BY COUNCIL DIRECTIVE 2007/47/EEC.

ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER.

NOTIFIED BODY: TÜV SÜD PRODUCT SERVICE GMBH

RIDLERSTR 65, D-80339 MUNCHEN, GERMANY

IDENTIFICATION NUMBER 0123

(EC) CERTIFICATE(S): No.G2 002145 0001 Rev.01

EC REP

EUROPEAN REPRESENTATIVE: MedNet EC-REP GmbH, Borstrasse 10, 48163. Muenster, Germany.

START OF CE-MARKING: 26/04/2019

PLACE, DATE OF DECLARATION: Shenzhen, 29/04/2021

SIGNATURE: XIACHANGCHUN

POSITION: GENERAL MANAGER

MANUFACTURER IS EXCLUSIVELY RESPONSIBLE FOR THE DOC.

Ref: EN ISO/IEC 17050-1 revision date: June 2009