## DECLARATION OF CONFORMITY TO COUNCIL DIRECTIVE 93/42/EEC CONCERNING MEDICAL DEVICES

MANUFACTURER: ZHEJIANG KINDLY MEDICAL DEVICES CO.LTD.

No.758, 5th Binhai Road, Binhai Industrial Park, Longwan District, 325025 Wenzhou, Zhejiang Province, PRC.

EUROPEAN REPRESENTATIVE: Shanghai International Holding Corp. GmbH (Europe)

Eiffestraße 80, 20537 Hamburg GERMANY

MEDICAL DEVICE: Scalp Vein Sets: 27G、26G、25G、24G、23G、22G、21G、

20G、19G、18G

CLASSIFICATION - ANNEX IX: CLASS IIA, RULE 7

CONFORMITY ASSESSMENT ROUTE: ANNEX II.3, Excluding(4)

We, <u>THE MANUFACTURER</u>, 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.

THE MANUFACTURER IS EXCLUSIVELY RESPONSIBLE FOR THE DECLARTION OF CONFORMITY

NOTIFIED BODY: TÜV SÜD PRODUCT SERVICE GMBH

RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY

IDENTIFICATION NUMBER C € 0123

(EC) CERTIFICATE(S): G1 036336 0054 Rev.02

START OF CE-MARKING: 2000.02

Valid until: 2024-05-26

PLACE, DATE OF DECLARATION: Wenzhou 2019-08-16

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SIGNATURE:

POSITION: QUALITY MANAGER