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

MANUFACTURER:

Edan Instruments, Inc.

#15 Jinhui Road, Jinsha Community, Kengzi Sub-District, Pingshan District, 518122 Shenzhen.

P.R.China

EUROPEAN REPRESENTATIVE:

Shanghai International Holding Corp. GmbH (Europe)

Eiffestrasse 80 D-20537 Hamburg Germany

PRODUCT/MODEL:

Digital Ultrasonic Diagnostic Imaging System/ DUS 60

The accessories are used together with the product

GMDN [NAME/CODE]:

General ultrasound imaging system, line-powered / 40761

CLASSIFICATION:

Class II a, Rule 10 According To Annex IX of the MDD

CONFORMITY ASSESSMENT ROUTE: Annex II excluding (4)

WE, EDAN INSTRUMENTS, INC., HEREWITH DECLARE THAT THE ABOVE MENTIONED PRODUCT(S) MEET THE TRANSPOSITION INTO NATIONAL LAW, THE PROVISIONS OF COUNCIL DIRECTIVE 93/42/EEC of 14 June 1993 including amendments by derective 2007/47/EC. ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER.

STANDARDS APPLIED: EN 60601-1:2006/A1:2013, EN 60601-1-2: 2007/AC:2010. EN 60601-2-37:2008, EN62366:2008, EN 62304:2006, EN ISO14971:2012, EN ISO 10993-1:2009, EN ISO10993-5:2009, EN ISO10993-10:2010, EN 980:2008, EN 1041: 2008

NOTIFIED BODY:

TÜV SÜD PRODUCT SERVICE GMBH

RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY

IDENTIFICATION NUMBER

CEO123

(EC) CERTIFICATE(S):

G1 16 04 91264 002 VALID UNTIL: 2017-09-17

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SIGNATURE:

MANAGEMENT REPRESENTATIVE