

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

MANUFACTURER:



TANITA CORPORATION
1-14-2, MAENO-CHO, ITABASHI-KU, TOKYO, JAPAN

EUROPEAN REPRESENTATIVE:



TANITA EUROPE B.V.
HOOGOORDDREEF 56-E
1101BE AMSTERDAM, THE NETHERLANDS

PRODUCT:

MC-780MA

SIMILAR MODEL:

NONE

UMDNS CODE:

17417

CLASSIFICATION:

CLASS IIA, (RULE 10)
RULE ACCORDING TO ANNEX IX OF THE MDD

CONFORMITY ASSESSMENT ROUTE: ANNEX II

WE UNDER OUR SOLE RESPONSIBILITY, HEREWITH DECLARE THAT THE ABOVE MENTIONED PRODUCTS MEET THE PROVISION OF THE COUNCIL DIRECTIVE 93/42/EEC CONCERNING MEDICAL DEVICES. A STATEMENT THAT THE MANUFACTURER IS EXCLUSIVELY RESPONSIBLE FOR THE DECLARATION OF CONFORMITY.

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

STANDARDS APPLIED:

SEE ATTACHED LIST OF "HARMONIZED STANDARD LIST"

NOTIFIED BODY:

TÜV SÜD PRODUCT SERVICE GMBH
RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY

IDENTIFICATION NUMBER



0123

START OF CE-MARKING:

THE APPLIANCE IS CE-MARKED 2013
SERIAL No. 13020001 –
(E.G. 13020001 = THE 1ST UNIT THAT HAS BEEN
MANUFACTURED FEB, 2013

PLACE, DATE OF ISSUE:

JAPAN, 1-JUN, 2017.

SIGNATURE:

DIRECTOR

BUSINESS STRATEGY H.Q. / LIFE SOLUTION DEPARTMENT