

EC Declaration of Conformity

Manufacturer: OMRON HEALTHCARE Co., Ltd.
Address: 53, Kunotsubo, Terado-cho, Muko, Kyoto 617-0002 JAPAN
European Representative: OMRON HEALTHCARE EUROPE B.V.
Address: Scorpius 33, 2132 LR Hoofddorp, The Netherlands
Product Category: Electronic Sphygmomanometers/Blood Pressure Monitors
Model Name(-code): M6 AC(HEM-7322-E)
Classification: Class IIa(MDD Annex IX Rule 10)

We herewith declare that the above mentioned product meets the provisions of the following European Committee Council Directives and Standards. All supporting documentation are retained under the premises of the manufacturer and the notified body.
This Declaration of Conformity is valid in connection with the shipping inspection reports for the respective batch of produced devices.

Directives

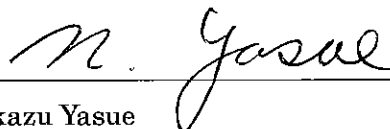
General applicable directives: Medical Device Directive (MDD) 93/42/EEC

Standards:
EN980:2008
EN1041:2008
EN1060-1:1995+A2:2009
EN1060-3:1997+A2:2009
EN60601-1:2006
EN60601-1-2:2007
IEC60601-1-6:2010
EN ISO14971:2012
EN ISO10993-1:2009
EN ISO10993-5:2009
EN ISO10993-10:2010
EN62304:2006
EN62366:2008
EN60601-1-11:2010
EN80601-2-30:2010

Notified Body: TÜV Rheinland LGA Products GmbH
Address: Tillystrasse 2, 90431 Nuremberg, Germany
ID No: Notified under number 0197 to the EC Commission
Certificate Registration No: Annex II : HD 60041154 0001

Place / Date: Kyoto / December 16, 2013

Signature:



Name: Norikazu Yasue
Position: General Manager
Customer Satisfaction Management Division