DECLARATION OF CONFORMITY

We,

ZOLL Medical Corporation 269 Mill Road Chelmsford, MA 01824-4105 USA,

declare under our sole responsibility that the products:

E Series E Series Pulse Oximetry Option E Series End Tidal CO₂ Option E Series 12 Lead Option E Series NIBP Option M Series M Series CCT M Series Biphasic Option M Series Pulse Oximetry Option M Series End Tidal CO₂ Option M Series 12 Lead Option M Series NIBP Option M Series IBP/TEMP Option AED PLUS AED PRO $CPR-D \ padz^{TM}$ R Series

to which this declaration relates are in conformity with the provisions of Council Directive 93/42/EEC of 14 June 1993 of the Medical Device Directive which apply to them.

The quality system under which these products were designed and manufactured has been certified by TUV Rheinland Product Safety (0197) to be in compliance with Annex II of the Medical Device Directive including European Standard, ISO 13485.

18 OCT 2006

Paul Días V.P. Quality Assurance and Regulatory Affairs

ZOLL Medical Corporation

Authorized European Representative Eric Rozeboom ZOLL International Holding B.V. Edisonring 3a 6669 NA Dodewaard The Netherlands Tel. +31 488-411-183

9652-0076 REV YA

ſF

Certificate of Conformance

This is to declare under our sole responsibility that the following ZOLL Medical Corporation products:

AED Plus AED Pro E Series M Series CCT Smart Battery Smart XL Battery R Series SurePower Battery

are in conformance with the provisions of Council Directive 2002/96/EC of 27 January 2003 on Waste Electrical and Electronic Equipment which apply to them.

