



Declaration of Conformity

Manufacturer: Philips Medical Systems
2301 Fifth Avenue, Suite 200
Seattle, WA 98121-1825
USA

European Representative: Philips Medizin Systeme Boeblingen GmbH
Hewlett-Packard Str. 2
71034 Boeblingen
Germany

Product: HeartStart HS1
Models – M5066A, M5067A, M5068A

Classification: Class IIb, Rule 9, Annex IX

Conformity Assessment Route: Annex II

We herewith declare that the above-mentioned products meet the provisions of the council Directive 93/42/EEC for Medical Devices. All supporting documentation is retained under the premises of the manufacturer.

Notified Body: TÜV SUD Product Service GmbH
Zertifizierstelle
Ridlerstrasse 65
D-80339 München
Germany

Start of CE-marking: October 9, 2007
Serial number A07J-01551

UMDNS Code: 17116/Defibrillators, Automatic, External

Place and Date of Issue: Seattle, WA / March 31, 2010

Signature Tom Trotter 3/31/2010
Tom Trotter, Sr. Manager, Regulatory Affairs Date