

**APPROVAL**  
EC Directive 93/42/EEC Annex II, Article 3  
Full Quality Assurance System  
Medical Devices

Registration No.: HD 60025225 0001

Report No.: 02421675 002

**Manufacturer:** HD MEDICAL SERVICES  
(INDIA) PRIVATE LIMITED  
No. 48, Industrial Estate, IT Highway  
Perungudi, Chennai 600096  
India

**Scope:** Design, development and manufacture  
of digital stethoscopes

**Date of Expiry:** 05.05.2014

The Notified Body hereby authorizes the quality management system established and applied by the company mentioned above. The requirements of Annex II, Article 3 of the directive have been met. This approval is subject to periodic surveillance, defined by Annex II, Article 5 of the aforementioned EC Directive, and can be used by the company with the manufacturer's declaration of conformity.

Notified Body



Cologne, 23.07.2009

Dipl.-Ing. D. Meier

**TÜV Rheinland Product Safety GmbH - Am Grauen Stein - D-51105 Köln**  
Accredited by Zentralstelle der Länder für Sicherheitstechnik (ZLS) and  
Zentralstelle der Länder für Gesundheitsschutz bei Arzneimitteln und Medizinprodukten (ZLG).

Notified under No. **0197** to the EC Commission.

CE The CE marking may be used if all relevant and effective EC Directives are complied with.

CE