## HITACHI

## 

MEDICAL SYSTEM OPERATIONS GROUP, KASHIWA 2-1 SHINTOYOFUTA, KASHIWA-SHI, CHIBA 277-0804, JAPAN

## EC-CONFORMITY DECLARATION

Manufacturer

Hitachi Medical Corporation

4-14-1, Soto-Kanda, Chiyoda-ku

Tokyo, 101-0021, Japan

European Representative

Hitachi Medical Systems GmbH

Otto von Guericke Ring 3,

D-65205 Wiesbaden, Germany

Medical Device

Modality

: Ultrasound Transducers / Probes

UMDNS Code

: 16-272 MDD Classification : IIa (Rule 5)

Model Name

: Transvaginal probe

Model Type

: EUP-VV531

The first manufactured serial number: Next to PE10654

This Declaration of Conformity covers all medical devices as specified in the product list belonging to this declaration and is only valid in connection with a batch specific Certificate of Compliance for all products concerned bearing the CE mark, as documented in each in process check list of the form No. SKM C2-125 for manufacturing.

The undersigned hereby declares that the medical device as specified above and related options comply with the essential requirements of Annex I of the EC-Directive 93/42/EEC.

The declaration of conformity is based on an assessment procedure in compliance with the EC Directive 93/42/ EEC, Annex II for a Full Quality Assurance System.

Notified Body: TÜV Rheinland LGA Products GmbH

Tillystraße 2, 90431 Nürnberg

Germany

Ident No.:

CE-0197

K. Kuriyama

General Manager

**Ultrasound System Division** 

N. Kawabe

**Division Manager** 

**Quality Assurance Division** 

Place **JAPAN** 

Date

2012-09-21