DECLARATION OF CONFORMITY

Manufactuer: Address:		Hitachi Aloka Medical, Ltd. 6-22-1 Mure, Mitaka-shi, Tokyo 181-8622 Japan
selected conformity assessment procedure:		Annex II excluding (4)
European Representative: Address:		Hitachi Medical Ultrasound Logistics, Zweigniederlassung der Hitachi Medical Systems Europe Holding AG Carl-Zeiss-Strasse 5 D-72555 Metzingen Germany
Product:		Transvaginal Electronic Convex Sector Scanner
Model Code::		ASU-1003
Classification (MDD, Annex IX):		II а
provisions	s of the following E	e above mentioned products including all its accessories meet the EC Council Directives and Standards. All supporting documentations hises of the manufacturer.
		DIRECTIVES
		Council Directive 93/42/EEC of 14 June 1993 concerning Medical Devices Directive (93/42/EEC as amended by 2007/47/EC; MDD)
Standards :	Harmonized Standards (published in the Office Journal of the European Communities) applicable to this product are :	
	EN 60601-1(2006)/AC(2010), EN 60601-1-2(2007)/AC(2010) EN 60601-2-37(2008), EN 62366(2008), EN 60601-1-6(2010) EN 980(2008), EN 1041(2008)	
Other Standa	ards :	
	EN ISO 14971(2009), EN ISO 10993-1(2009)/AC(2010)	

EN ISO 14971(2009), EN ISO 10993-1(2009)/AC(2010) IEC 62079(2001), ISO 7010(2003)/A1(2006) ISO 3864-2(2004)

Notified body :TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123Address:Ridlerstraße 65, 80339 München, Germany

Start of CE marking: SN : 202A6923

Place :

Hitachi Aloka Medical,Ltd. Tokyo Works

Date: **31. May. 2012**

Signature:

MASAAKI HIRAKAWA

Name of issuer :Masaaki HirakawaPosition :General Manager of Quality Assurance Department