

EC-DECLARATION OF CONFORMITY

Manufacturer: Address:	Hitachi, Ltd. 2-16-1 Higashi-Ueno, Taito-ku, Tokyo, 110-0015, Japan
Selected conformity assessment procedure: Annex II excluding (4) RoHS Article 7 (b), Module A	
EU Authorized represer Address:	ntative: Hitachi Medical Systems GmbH Otto-von-Guericke-Ring 3, D-65205 Wiesbaden, Germany
Product: Model Code <mark>REF</mark> :	S21 Probe
Classification (MDD, Annex IX): IIa Categories (RoHS(II), Annex I): No.8 Classification rule (MDD, Annex IX): Rule 10	
Statement: We are exclusively responsible for the declaration of conformity and herewith declare that the above-mentioned product including all its options meet the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer.	
DIRECTIVES	
General applicable directives: Medical Device Directive : Council Directive 93/42/EEC of 14 June 1993 as amended by 2007/47/EC;	
Notified body : Address (for MDD):	TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123 Ridlerstraße 65, 80339 München, Germany
RoHS Directive :	Directive 2011/65/EU of 8 June 2011 concerning on the restriction of the use of certain hazardous substances in electrical and electronic equipment.
Production facility : Address:	Hitachi Healthcare Manufacturing, Ltd. Analytical Systems Kashiwa Factory 2-1, Shintoyofuta, Kashiwa-shi, Chiba, 277-0804 JAPAN
Starting of CE Marking: Date:	<u>G3031267</u> Sep.03,2019

Signature:

Name of issuer : Position :

Anaeda Ryosuke Maeda

Management Representative

Place: Tokyo, JAPAN