

## **EC-DECLARATION OF CONFORMITY**

Manufacturer: Hitachi, Ltd.

Address: 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 representative: Hitachi Medical Systems GmbH

Address: Otto-von-Guericke-Ring 3, D-65205 Wiesbaden, Germany

Product: Transrectum Puncture Electronic Convex Probe

Model Code REF: UST-676P

Classification (MDD, Annex IX): IIa
Categories (RoHS(II), Annex I): No.8
Classification rule (MDD, Annex IX): Rule 5

## 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: TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

Address (for MDD): 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: Hitachi Healthcare Manufacturing, Ltd. Analytical Systems Kashiwa Factory

Address: 2-1, Shintoyofuta, Kashiwa-shi, Chiba, 277-0804 JAPAN

Starting of CE Marking: G3005035

Date: Jun.10,2019

Signature:

Name of issuer: Ryosuke Maeda Place: *Tokyo, JAPAN* 

Maeda

Position: Management Representative