

### Declaration of Conformity (No. E2020006)

We, Sun Medical Co., Ltd., ensure and declare that the products listed in the schedule A below meet the provisions of the Directive 93/42/EEC concerning medical devices. Our products listed in the schedule A below are also in conformity with the safety requirements laid down in Annex I of the Directive 93/42/EEC as amended and the declarations of compliance of our products listed in the schedule A are issued according to Annex II excl. section 4 of the Directive 93/42/EEC as amended on Medical Devices.

EC Certificate No.: G1 026583 0028


Our authorized representative in EU is: EMERGO EUROPE B.V. having its principal office at Prinsessegracht 20, 2514 AP, The Hague, The Netherlands.

The notified body is: TÜV SÜD Product Service GmbH (ID No. 0123) at Ridlerstrasse 65. 80339 Munich, Germany

#### Schedule A (Part 1)

Product	Type of Product	Classification (Applied Rule of Annex)	Lot Number From:	Standards Applied		Additional Information
Hybrid Bond ONE	Dental Bonding Agent	Ila (Annex IX, Rule 8)	In Part 2	EN ISO 15223-1:2016 EN 1041:2008 EN 1641:2009 EN ISO 7405:2008	EN ISO 10993-1:2009 EN ISO 10993-18:2009 EN ISO 14971:2012 EN 62366:2008	-

We are exclusively responsible for this statement.

 July, 2, 2020

Narimichi Honda

Management Representative for Quality Management System

Sun Medical Co., Ltd.

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**Schedule A** (Part 2)

<b>Product code</b>	<b>UDI-DI</b>	<b>Product name</b>	<b>Lot number From:</b>
T435E	4560227799782	Hybrid Bond ONE	TV1
T435TR	4560227800655		