

EC Declaration of Conformity

Manufacturers Name: Asiga

Manufacturers Address: Unit 2, 19-21 Bourke Rd, Alexandria, NSW, 2015, Australia

SRN (Single Registration Number): AU-MF-000012099

Authorized Representative Name

(if applicable):

MT Promedt Consulting GmbH

Authorized Representative

Address (if applicable):

Ernst-Heckel-Straße 7, 66386 St. Ingbert, Germany

SRN (Single Registration Number): DE-AR-000000085

Basic UDI-DI: 93592150RESINSUW

Name of the Device (s): DentaGUIDE, DentaTRAY, DentaTRY, DentaIBT

Product Code / Part Number: DentaGUIDE: 04504

DentaTRAY: 05165 DentaIBT: 03746

DentaTRY: A1: 03768, A2: 03817, A3: 03819,

B1: 03821, B2: 03823, B3: 03825

Classification: Class I per Rule 5 of Annex VIII of the Medical Devices Regulations (EU 2017/745)

Intended Purpose: The subject products are raw materials intended to be used for additive

manufacture in combination with Digital Light Processing (DLP) based 3D printers that support Asiga resins to manufacture parts for Dental devices.

Notified Body name: N/A – Class I, Self-Certified.

Notified Body Address: N/A

Notified Body Identification

number:

N/A

Common Specifications (CS): N/A

Conformity assessment route: Medical Device Regulation 2017/745, Article 52(7) MDR

This declaration of conformity is issued under the sole responsibility of Asiga. We hereby declare that the medical device(s) specified above meet the provision of the Regulation (EU) MDR 2017/745 for medical devices. This declaration is supported by the Quality System approval to ISO 13485 issued by BSI. All supporting documentation is retained at the premises of the manufacturer.

Signature: Place and date of issue:

Just Co-y Alexandria, Sydney, Australia, 30/05/2023

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Justin Elsey

Managing Director

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