

EU Declaration of Conformity

to REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE
COUNCIL of 5 April 2017 on medical devices

Manufacturer: Greiner Bio-One GmbH SRN: N/A
Bad Haller Straße 32
4550 Kremsmünster
Austria

Production Greiner Bio-One GmbH Greiner Bio-One (Thailand) Ltd.
Location: Bad Haller Straße 32 700/172 Moo.1
4550 Kremsmünster Amata Nakorn Industrial Estate
Austria Tambon Bankao
Amphur Phanthong
Chonburi 20160
Thailand

Product / Tube Holder
Product Group: (for details please refer to page 2)

Basic UDI-DI (GMN): (for details please refer to page 2)

Classification: Class I according to Regulation (EU) 2017/745 of the european parliament and of
the council of 5 April 2017 on medical devices, Annex VIII - Chapter III
Classification Rules - 4.1. Rule 1

GMDN Code(s): 37566

We herewith declare under our sole responsibility that the products specified above meet the provisions of the above-mentioned Regulation. All supporting documentation is retained under the premises of the manufacturer.


Conformity Assessment procedure according to Annex IV of the Regulation (EU) 2017/745.

Standards / common specifications:

Refer to the list of applicable (harmonized) standards and common specifications in the Technical Documentation.

Kremsmünster, 10.05.2021




Georg Sambs
Quality Manager Greiner Bio-One Austria



Product Group	Product name - detailed product description	Item numbers	Basic UDI-DI (GMN)
Tube Holder	Standard Tube Holder 10 pcs. per bag	450201	912001757G000000199
Tube Holder	Standard Tube Holder 100 pcs. per bag	450209	912001757G000000199
Tube Holder	Speedy Quick Release Holder	450212	912001757G00000039D
Tube Holder	VACUETTE® QUICKSHIELD Safety Tube Holder	450230	912001757G00000029B