

EU Declaration of Conformity

to REGULATION (EU) 2017/746 OF THE EUROPEAN PARLIAMENT AND OF THE
COUNCIL of 5 April 2017 on in vitro diagnostic medical devices

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

Production Greiner Bio-One GmbH
Location: Bad Haller Straße 32
4550 Kremsmünster
Austria

Product / MiniCollect® K2E K2EDTA TUBE
Product Group: (for details please refer to page 2)

BASIC-UDI-DI (GMN): 912001757G00000439R

Classification: Class A according to Regulation (EU) 2017/746 of the european parliament and of
the council of 5 April 2017 on in vitro diagnostic medical devices, Annex VIII
Classification Rules - Rule 5

GMDN Code(s): 58144

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 acc. to Annex IV of the Regulation (EU) 2017/746.

Standards / common specifications:

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

Kremsmünster, 20.05.2022


Georg Sambs
Quality Manager
Greiner Bio-One Austria

PRODUCT GROUP	Product name - detailed product description	Item numbers
MiniCollect® K2E K2EDTA TUBE	MiniCollect® TUBE 0.25 / 0.5 ml K2E K2EDTA lavender cap	450532
MiniCollect® K2E K2EDTA TUBE	MiniCollect® Complete 0.25 / 0.5 ml K2E K2EDTA lavender cap, pre-assembled with Carrier Tube 13x75	450547
MiniCollect® K2E K2EDTA TUBE	MiniCollect® Complete 0.25 / 0.5 ml K2E K2EDTA lavender cap, paper label, pre-assembled with Carrier Tube 13x75	450647