

EU Declaration of Conformity

In accordance with the EU Low Voltage Directive 2014/35/EU, Annex IV

We,

Trotec GmbH

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declare in sole responsibility that the following device

Product model / Product: BO 21
Year of manufacture as of: 2014
Product, batch, type or serial number: endoscope

has been developed, constructed and manufactured in compliance with the requirements of the European Directives:

2014/30/EU EMV Guideline
2011/65/EU RoHS 2 Guideline
2014/35/EU low voltage directive
2015/863/EU RoHS Guideline Annex II Phthalate complement
Guideline 2012/19/EU WEEE2- Guideline

The object of the declaration described above is in conformity with the relevant Union harmonisation legislation.

The assessment is based on the following applied harmonised standards:

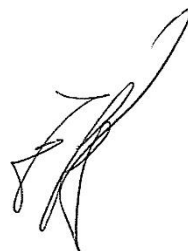
EN 61326-1:2013
EN 61326-2-2:2013
EN ISO 12100:2010

Other applied technical standards and specifications:

IEC 61000-4-2:2008
IEC 61000-4-3:2010
IEC 61000-4-8:2009
abgel. CISPR 11:2009/AMD1:2010-03
Regulation (EG) Nr. 1907/2006 (REACH)
Regulation (EG) Nr. 552/2009 (REACH Annex XVII)
Regulation (EU) 2018/2005 amending the Annex XVII
REACH

Place and date of issue:

Heinsberg, 28/09/2021



Joachim Ludwig, Managing Director

 **TROTEC**

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