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## **EC Declaration of Conformity**

We, TaiDoc Technology Corporation

B1-7F, No.127, Wugong 2nd Road, Wugu Dist., 24888 New Taipei City, TAIWAN declare under our sole responsibility that the product

**Product Name** 

: Sterile Blood Lancet

**Product Model** 

: TD-5084

Classification

: 93/42/EEC(Directive including 2007/47/EC)(MDD),

Annex IX, Section 2, Rule 6, Class IIa

Conformity Assessment Route

: 93/42/EEC(Directive including 2007/47/EC)(MDD)

Annex II excluding (4)

EC Certificate Number

: G1 052126 0043 Rev.03

European Representative

: MedNet EC-REP GmbH

Borkstraße 10, 48163 Münster, Germany

Notified Body (CE0123)

: TÜV SÜD Product Service GmbH

Ridlerstraße 65, 80339 München, Germany

GMDN code

: 45142

## to which this declaration relates is in conformity with the following standard(s) or other normative document(s):

ISO 10993-1:2018	Biological evaluation of medical devices Part 1: Evaluation and testing within a risk management process
	testing within a risk management process
ISO 10993-5:2009	Biological evaluation of medical devices. Tests for in vitro
	cytotoxicity
ISO 10993-10:2010	Biological evaluation of medical devices. Tests for irritation and
	skin sensitization
ISO 10993-12:2012	Biological evaluation of medical devices. Sample preparation
	and reference materials
EN ISO 11137-1:2015	Sterilization of health care products - Radiation - Part 1:
	Requirements for development, validation and routine control of
	a sterilization process for medical devices
EN ISO 11137-2:2015	Sterilization of health care products - Radiation - Part 2:
	Establishing the sterilization dose



## 泰博科技股份有限公司 TaiDoc Technology Corp.

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www.taidoc.com	<b>"</b> "是一个人们的一个人们的一个人们的一个人们的一个人们的一个人们的一个人们的一个人们的
EN ISO 11607-1:2009+A1:2014	Packaging for terminally sterilized medical devices - Part 1:
	Requirements for materials, sterile barrier systems and packaging
	systems
EN ISO 11607-2:2006+A1:2014	Packaging for terminally sterilized medical devices - Part 2:
	Validation requirements for forming, sealing and assembly
	processes
EN ISO 11737-1:2018	Sterilization of medical devices - Microbiological methods - Part1:
	Determination of population of microorganisms on products.
EN ISO 11737-2:2009	Sterilization of medical devices - Microbiological methods - Part 2:
	Tests of sterility performed in the definition, validation and
	maintenance of a sterilization process
ISO 13485:2016	Medical devices - Quality management systems - Requirements for
	regulatory purposes.
EN ISO 14971:2012	Medical devices - Application of risk management to medical
	devices.
EN ISO 15223-1:2016	Medical devices - Symbols to be used with medical device labels,
	labelling, and information to be supplied - Part 1:General
	requirements
EN 556-1:2001/AC: 2006	Sterilization of medical devices - Requirements for medical devices
	to be designated "STERILE" - Part 1: Requirements for terminally
	sterilized medical devices
EN 556-2:2015	Sterilization of medical devices - Requirements for medical devices
	to be designated "STERILE" - Part 2: Requirements for aseptically
	processed medical devices
EN 62366-1:2015	Medical devices - Application of usability engineering to medical
	devices

2021.5,13

Date of Issue

Jim Van

Management Representative Jim Jan