## URIT优利特

## **CE Technical Document**

**Declaration of Conformity** 

(Sheath)

(Model:URIT 5S 11)

Document No: IVDR-URIT 5S 11-CE09

Version/Revision No: 1.1

Department Distributed: Department II

Prepared by:	Verified by:	Approved by:
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Technical Engineer	Technical Manager	PRRC

## **Declaration of Conformity**

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	URIT Medical Electronic Co., Ltd.			
Manufacturer:	Address: No. D-07 Information Industry			
	District, High-Tech Zone, Guilin, Guangxi			
	541004, P. R. China			
	Shanghai International Holding Corp. GmbH			
European Representative:	(Europe)			
European representative.	Address: Eiffestrasse 80, 20537 Hamburg,			
	Germany			
Product Name:	Sheath			
Model:	URIT 5S 11			
GMDN Code:	61165			
EMDN Code:	W0103010105			
SRN:	CN-MF-000011840			
Basic UDI-DI:	69357404IBHS000006BB			
Device Photograph:	USIT STATE OF THE PARTY OF THE			
	It is used to dilute blood samples to form sheath flow, which is conducive to cell			
Intended Use:	counting and classification by analytical			
	instruments.			
Risk Class:	Class A			
	Rule 5 (a) of Annex VIII of the Regulation			
Classification Rule:	(EU) 2017/746			
	Article 48 section 10 of the Regulation (EU)			
Conformity Assessment Route	2017/746			
We herewith declare that the above-mentioned product(s) meet the Regulation (EU) 2017/746 of				
THE EUROPEAN PARLIAMENT AND OF TH	E COUNCIL. All supporting documentation is			
retained at the premises of the manufacturer. We, the manufacturer, are exclusively responsible for				
the DoC.				
	EN ISO 13485:2016			
Applied Standards:	EN ISO 14971:2019			
	EN ISO 15223-1:2021			
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		EN ISO 18113-2: 2022	
		EN ISO 18113-2: 2022 EN ISO 18113-1:2022	
		EN ISO 23640: 2015	
		EN 13612: 2002	
Matified Dodge		NA	
Notified Body:			
Identification Number:		NA	
(EC) Certificate(s):	NA	Valid until	NA
Start of CE-marking:			
Place, Date of Issue:		Guilin, Guangxi 2023-12-8	
Signature:		Name: Shi Ping Function: CEO	
		35030960V56b	