E 60160016	Document name	Declaration of Conformity	Document No.:	GD/MDR 10-08	
	Product name	Disposable Plastic Forceps	Version	1.0	Page 1 of 1

Declaration of Conformity

Manufacturer:	Name and Address
	Name:Zhejiang Gongdong Medical Technology Co., Ltd.
	Registered address:
	No.10 Beiyuan Ave., Huangyan, 318020 Taizhou, Zhejiang, People's
	Republic of China.
	Production address:
	(1) No.10 Beiyuan Ave., Huangyan, 318020 Taizhou, Zhejiang,
	People's Republic of China.
	(2) No.39 Beiyuan Ave., Huangyan, 318020 Taizhou, Zhejiang,
	People's Republic of China.
	(3) No.88 Jingxian Road, Huangyan, 318020 Taizhou, Zhejiang,
	People's Republic of China.
SRN of the Manufacturer:	CN-MF-000005694
Authorised Representative:	Shanghai International Holding Corp. GmbH(Europe)
	Eiffestrasse 80, 20537 Hamburg, Germany
SRN of the Authorised Rep.:	DE-AR-00000001
Product Name:	Disposable Plastic Forceps
Basic UDI-DI of Product:	6947462411111192M8
Intended Purpose:	The product is intended to be used to for collection, transportation
	and storage of samples.
EMDN Code:	Q030304

Classification (MDR, Annex VIII): I, rule 1

Conformity Assessment Procedure: ursuant to Regulation(EU)2017/745 on Medical Devices, Annex IX Chapters I and III.

We (manufacturer) herewith state that the above-mentioned product is in conformity with the following In Vitro Diagnostic Regulation, Common Specifications and Product Standards. We are solely responsible for the EU declaration of conformity.

The applicable In Vitro Diagnostic Regulation, Common Specifications and Product Standards: Medical

Device Regulation (EU) 2017/745

Reference Standards: EN ISO 14971: 2019 EN ISO 20417:2021 EN ISO 13485:2016 Signature:

Name: Position: Place, Date of Issue: EN ISO 18113-1: 2011 EN ISO15223-1:2021

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Zhong Weifeng General Manager Tai Zhou, 2023.10.07