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EC DECLARATION OF CONFORMITY

Durrani & Co.

We,

Naul, Wazirabad Road, Sialkot-51310-Pakistan

Hereby under sole responsibility declares that below mentioned medical devices manufactured by us have been classified according to the classification rules stated in the Annex – IX and conform to the ESSENTIAL REQURIEMENT as laid out in MDD 93/42/EEC in Annex-I and change of classification will be updated till 31-Dec-2028 according to the EU MDR 2017/745 transition timeline as amended by 2020/561 and the CE marking may be affixed.

Device Name:

Intended Purpose:

A hand-held manual instrument designed primarily for non-dedicated grasping of devices, sponges and/or dressings during a procedure; it is neither intended for tissue manipulation nor endoscopic use.

Annex 1

Device Classification:	Class: I	Rule: 6	Indent:	2
Single Registration Number (SRN):		PK-MF-000016079		

Product List: Annex 1

Sr. #	Product Code / Catalogue #	Product name
1	04-010-140	Dressing Forceps, Straight, No Teeth, 14 cm

Note:

Refer to respective Summary of Technical Documentation (STED), for product's photograph.



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Reference Regulation(s) / Standard(s) / Guidance Document(s) / Common Specification(s) (CS):

To which this declaration related is in conformity with the following standard(s) or other normative document(s)

Sr.	Description	Standards/Regulation/CS
	Medical devices – Quality management systems-Requirements for	
1	regulatory purposes.	EN ISO 13485:2016+AC:2018
	Medical Device Regulation (EU) of the European Parliament and of	EU MDR 2017/745 as
2	the Council	amended by 2020/561
3	Guidance notes for manufacturers of class I medical devices	MDCG 2019-15 rev.1
4	Conformity assessment – Supplier's declaration of conformity – Part 1: General requirements	EN ISO/IEC 17050-1:2010
		EN 150/12010
5	Medical devices-Application of risk management to medical Devices	EN ISO 14971:2019
	Medical Devices – Symbols to be used with medical device labels,	
	labeling and information to be supplied part-1 –general	
6	requirement	EN ISO 15223-1:2016
	Medical devices – Symbols to be used with medical device labels,	
	labeling, and information to be supplied – Part 2: Symbol	
7	development, selection and validation	ISO 15223-2:2010
	Biological Evaluation of medical Devices – Part 1: Evaluation and	
8	testing within a risk management process	EN ISO 10993-1:2020
	Biological evaluation of medical devices – Part 5: Tests for in vitro	
9	cytotoxicity	EN ISO 10993-5:2009
10	Biological evaluation of medical devices – Part 10: Tests for irritation	EN ISO 10993-10:2013
11	Standard Specification for Wrought Stainless Steel for Surgical	ASTM F899-20
12	Surgical Instruments – Metallic Materials – Part 1: Stainless Steel	EN ISO 7153-1:2016
13	Surgical and Dental Hand Instruments – Determination of resistance	EN ISO 13402:2000
14	Standard Test Method for Corrosion of Surgical Instruments	ASTM F1089-18
15	Clinical evaluation – guide for manufacturer and notified bodies	MEDDEV 2.7/1 Rev.4
16	Guidance on clinical evaluation – Equivalence	MDCG 2020-5
17	Guidance on sufficient clinical evidence for legacy devices	MDCG 2020-6
18	Best Practices for Medical Device Post Market Surveillance	ISO/TR 20416:2020
19	Post-marketing surveillance (PMS) Recommendation	NB-Med 2_12-1_rev11
20	Guidance on a Medical devices vigilance system	MEDDEV 2.12/1 rev.08
21	Processing of health care products – Information to be provided by	EN ISO 17664:2017
22	Sterilization of Health care products – Moist Heat – Part 1:	EN ISO 17665-1:2006



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Sr.	Description	Standards/Regulation/CS	
23	Comprehensive guide to steam sterilization and sterility assurance	ANSI / AAMI ST79:2017	
24	Medical devices – Part 1: Application of usability engineering to	EN 62366-1:2015/A1:2020	
25	Safety aspects — Guidelines for the inclusion in standards	ISO/IEC Guide 51:2014	
26	Guide to the development and inclusion of aspects of safety in	ISO/IEC GUIDE 63:2019	

EU AUTHORIZED REPRESENTATIVE (EUAR):

CMC MEDICAL DEVICES & DRUGS, S.L C/ Horacio Lengon18 C. P 29006 Málaga-Spain

Phone: +34 951 214 054 Email: info@cmcmedicaldevices.com

Signed for and on behalf of:

Name:	Gohar Durrani
Designation:	Head of Regulatory Compliance
Place of Issue:	Naul, Wazirabad Road, Sialkot-51310-Pakistan
Date of Issue:	8-Apr-21



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