MY TICARET VE MEDIKAL A.S.



Ömerli Mahallesi General Şükrü Koraltı Caddesi No:33 Arnavutkoy –Istanbul Turkey Tel: +902124382064 Fax: +902124382065 Website: <u>www.mymedikal.com.tr</u>.

## EU DECLARATION OF CONFORMITY

We, MY TICARET VE MEDIKAL A.S. herewith declare under our sole responsibility that below mentioned product{s) with CE mark is fully compliance with Essential Requirement of the EC Council Directive 93/42/EEC 14th June 1993 concerning medical devices, amended by Council Directive 2007/47/EC.

EC Certificate	Not applicable (Self- declared)			
Manufacturer	MY TICARET VE MEDIKAL A.S.			
Manufacturer Address	Ömerli mah General Şükrü Koraltı Cd No:33, 34555			
		utkoy/Istanbul, Turkey	-	
Single Registration Number	TR-MF-000018372			
(SRN)				
Brand	Mumu Plus+			
Product Description	Powderfree Nitrile Gloves			
Intended Purpose	A patient examination glove is a medical device intended for a medical purpose that is worn on the examiners hand or finger to prevent contamination between patient and examiner. Examination glove is intended for medical activities except surgery.			
Size	XS, S, M, L, XL			
Colors	Blue, Cool Blue, Black			
European Medical Device	T01020204 (Examination / Treatment Gloves, Nitrile)			
Nomenclature (EMDN)				
Global Medical Device	56286 (Nitrile Examination/Treatment glove, non-			
Nomenclature (GMDN)	powdered, non-sterile)			
Product Catalogue Number	MN01			
Conformity Assessment Route	Annex VII			
Classification & Rule	Class I, Rule 5			
Applicable Standards				
	No.	Regulation/ Standard Number	Regulation/ Standard Name	
	1	MDD 93/42/EEC	Medical Device Directive	
			Medical devices - Quality	
	2	ISO 13485: 2016	management systems -	
			Requirements for regulatory purposes	
			Quality management systems –	
	3	ISO 9001: 2015	requirements	
	4	ISO 14971: 2019	Medical devices - application of risk	



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			management to medical devices
	5	EN 455-1: 2020	Requirements and testing for
			freedom from holes
	6	EN 455-2: 2015	Requirements and testing for
			physical properties
	7	EN 455-3: 2015	Requirements and testing for
			biological evaluation
	8	EN 455-4: 2009	Requirements and testing for
			shelf-life determination
	9	ISO 10993-10: 2010	Biological evaluation of medical
			devices – Part 10: Test for irritation
			and skin sensitization
	10	EN 1041: 2008+A1: 2013	Information supplied by the
			manufacturer of medical devices
		11 ISO 15223-1: 2021	ISO 15223-1 Symbols to be used
	11		with information to be supplied
			by the manufacturer

The gloves are manufactured according to EN ISO 9001:2015 and EN ISO 13485:2016 Quality Management System:

Verification Certificates: EN ISO 13486:2016 Quality Management System Certificate No: ISO 02 836 1179

> EN ISO 9001:2015 Quality Management System Certificate No.: ISO 01 940 1179

## Authorized Signatory:

Approver	: MURAT YILDIZ
Title	: General Manager/CEO
Signature	No:33 AMAYUKÖY/ISTANBUL
Approval Date	el:02174182064Fax:02124382065 : 26.10,2022
Place of Approval	: Istanbul, Turkey

## CE