

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

EC Certificate	Not applicable (Self- declared)			
Manufacturer	MY TICARET VE MEDIKAL A.S.			
Manufacturer Address	Öme	rli mah General Şükrü K	oraltı Cd no:33, 34555	
	Arna	vutkoy/Istanbul, Turkey	,	
Single Registration Number	TR-M	F-000018372		
(SRN)				
Brand	Mumu Plus+			
Product Description	Powdered Latex Examination Gloves			
Intended Purpose	A patient examination glove is a medical device intended for			
			vorn on the examiners hand or	
			ination between patient and	
	_		ove is intended for medical	
	activities except surgery.			
Basic UDI-DI	868227994LP5H			
Size	XS, S, M, L, XL			
European Medical Device	T010	T010201 (Examination/Treatment Gloves, Latex)		
Nomenclature (EMDN)		•		
Global Medical Device	4717	47173 (Latex examination/treatment glove, powdered)		
Nomenclature (GMDN)				
Product Catalogue Number	SMLP00-XS; SMLP01-S; SMLP02-M; SMLP03-L; SMLP04-XL			
Conformity Assessment Route	Annex II and Annex III according to EU 2017/745			
(MDR):				
Classification & Rule (MDR)	Class I, Rule 5 transient use according to Annex VIII			
Device Classification (PPER)	Category III			
Product Group Reference	SSLPLP			
Number				
EU Type-Examination	2777/12636-01/E03-02			
Certificate (PPER)				
Notified Body Number (PPER)	2777	2777		
EU Type- Examination	SATR	A Technology Europe Li	imited	
Certificate Issued by (PPER)		• · · ·		
Applicable Standards	Bracetown Business Park, Clonee, D15YN2P, Ireland			
	No.	Regulation/ Standard Number	Regulation/ Standard Name	
	1	MDR (EU) 2017/745	Medical Device Regulation	
			Personal Protective Equipment	
	2	PPE (EU) 2016/425	Regulation	
	3	ISO 13485: 2016	Medical devices - Quality management	
	1.1			



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			systems - Requirements for
			regulatory purposes
			Quality management systems –
	4	ISO 9001: 2015	requirements
			Medical devices - application of risk
	5	ISO 14971: 2019	management to medical devices
	6	EN 455-1: 2020	Requirements and testing for
			freedom from holes
	7		Requirements and testing for
		EN 455-2: 2015	physical properties
	8	EN 455-3: 2015	Requirements and testing for
			biological evaluation
		EN 455-4: 2009	Requirements and testing for
	9		shelf-life determination
			Biological evaluation of medical
	10	ISO 10993-10: 2010	devices –Part 10: Test for irritation
	10	150 10555 10. 2010	and skin sensitization
			Information supplied by the
	11	EN 1041: 2008+A1: 2013	manufacturer of medical devices
			ISO 15223-1 Symbols to be used
	12	ISO 15223-1: 2021	with information to be supplied
			by the manufacturer
			Protective gloves against
	13	EN ISO 374-1: 2016	u
			dangerous chemicals and micro-
			organisms - Part 1: Terminology
			and performance requirements for chemical risks
	14	EN ISO 374-2:2014	Protective gloves against
			dangerous chemicals and micro-
			organisms - Part 2: Determination
			of resistance to penetration
		EN ISO 374-4:2013	Protective gloves against chemicals
	15		and micro-organisms - Part 4:
			Determination of resistance to
			degradation by chemicals
			Protective gloves against
			dangerous chemicals and micro-
	16	EN ISO 374-5: 2016	organisms - Part 5: Terminology
			and performance requirements for
			micro-organisms risks
	17	EN 16523-1: 2015	Determination of material
			resistance to permeation by
		LIN 10323-1. 2013	chemicals - Part 1: Permeation by
			liquid chemical under conditions of
			continuous contact



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We, My Ticaret ve Medikal A.S. herewith declare that the above-mentioned device:

- Is in compliance with the General Safety Performance Requirement of the Medical Device Regulation (MDR) 2017/745. All supporting documentations are retained under the premise of 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

- Is following to the EU-Type Examination and conformity with the provisions of new PPE Regulations (EU) 2016/425 Category III of the notified body number 2777 by SATRA Technology Europe Ltd.
- Is in conformity to type based on quality assurance of the production process under surveillance of the notified body number 2777 by SATRA Technology Europe Ltd.
- This EU Declaration of Conformity is prepared in accordance to Annex IV of Medical Device Regulation (EU) 2017/745.

Authorized Signatory:

Approver	: MURAT YILDIZ
Title	: General Manager/CEO
Signature	MY ICARETVE MEDIKALIAN ONIM SIRKETI Omeril Mah Septin Sürke Ti No:33 AM VKöyilSTANBUL Büyüka MUKA VKÖyilSTANBUL Büyüka MUKA VL 0.826 040 4605
Approval Date	et:021743920 64 Fax:0212 438 20 65 : 25-10 2022 ethel.com
Place of Approval	: Istanbul, Turkey

