

MY TICARET VE MEDIKAL A.S.

Ömerli Mahallesi General Şükrü Koraltı Caddesi No:33 Arnavutkoy –Istanbul Turkey Tel: +902124382064 Fax: +902124382065

Website: www.mymedikal.com.tr.

EU DECLARATION OF CONFORMITY

DOC No.	DOC-MYMEDIKAL-SRT-003				
EC Certificate	Not applicable (Self- declared)				
Manufacturer	MY TICARET VE MEDIKAL A.S.				
Manufacturer Address	Ömerli mah General Şükrü Koraltı Cd No:33, 34555				
	Arnavutkoy/Istanbul, Turkey				
Single Registration Number (SRN)	TR-MF-00018372				
Brand	B-good				
Product Description	Latex Powderfree Examination Gloves				
Intended Purpose	A patient examination glove is a medical device intended for a medical purpose that is worn on the examiner's hand or finger to prevent contamination between the patient and examiner. Examination glove is intended for medical activities except for surgery.				
Basic UDI-DI	868227994LPFX3				
Size	XS, S, M, L, XL				
European Medical Device Nomenclature (EMDN)	T010201 (Examination/Treatment Gloves, Latex)				
Global Medical Device	47172 (Hevea-latex Examination/treatment glove, non-				
Nomenclature (GMDN)	powdered, non-antimicrobial)				
Product Catalogue/Reference Number	BGLP01-XS, BGLP02-S, BGLP03-M, BGLP04-L, BGLP05-XL				
Product Group Reference	LO01				
Number					
Conformity Assessment Route (MDR):	Annex II and Annex III according to EU 2017/745				
Classification & Rule (MDR)	Class I, Rule 1 & Rule 5				
Device Classification (PPER)	Category III				
EU Type-Examination Certificate (PPER)	2777/10467-05/E15-01				
Notified Body (PPER)	EU-Type Examination and Ongoing Conformity by Notified Body SATRA TECHNOLOGY EUROPE LTD				
	Bracetown Business Park,				
	Clonee, D15YN2P, Ireland [CE 2777]				
Applicable Standards	cionec, 515 (142) , il ciuna [et 2777]				
Applicable Statidatus	No. Regulation/ Standard Regulation/ Standard Name Number				
	1 MDR (EU) 2017/745 Medical Device Regulation				
	2 PPE (EU) 2016/425 Personal Protective Equipment Regulation				
	3 ISO 13485: 2016 Medical devices - Quality management				

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		systems - Requirements for
		regulatory purposes
4	ISO 9001: 2015	Quality management systems – requirements
5	ISO 14971: 2019	Medical devices - application of risk
		management to medical devices
6	EN 455-1: 2020	Requirements and testing for
		freedom from holes
7	EN 455-2: 2015	Requirements and testing for
		physical properties
8	EN 455-3: 2015	Requirements and testing for
		biological evaluation
9	EN 455-4: 2009	Requirements and testing for
		shelf-life determination
10	ISO 10993-1: 2018	Biological evaluation of medical
		devices –Part 1: Evaluation and
		testing within a risk management
		process
11	ISO 10993-5: 2009	Biological evaluation of medical
		devices –Part 5: Test for in vitro
		cytotoxicity
12	ISO 10993-10: 2010	Biological evaluation of medical
		devices –Part 10: Test for irritation
- 10	ACTA 54674 0040	and skin sensitization
13	ASTM F1671: 2013	Standard test method for resistance
		of materials used in protective
		clothing to penetration by blood-
		borne pathogens using phi-x174 bacteriophage penetration as a test
14	ASTM D3578: 2019	system Standard specification for rubber
17	A3110 D3370. 2013	examination gloves
15	ISO 20417:2021	Medical devices - Information to be
	130 20717.2021	supplied by the manufacturer
16	ISO 15223-1: 2021	ISO 15223-1 Symbols to be used
		with information to be supplied
		by the manufacturer
17	ASTM D7160: 2016	Determination of expiration
		dating for medical gloves
18	ASTM D7161: 2016	Determination of real time
		expiration dating of mature
		medical gloves stored under typical
		warehouse conditions
19	EN ISO 374-1: 2016+A1:	Protective gloves against
	2018	dangerous chemicals and micro-
		organisms - Part 1: Terminology
		and performance requirements for
		chemical risks

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20	EN ISO 374-2: 2019	Protective gloves against
		dangerous chemicals and micro-
		organisms - Part 2: Determination
		of resistance to penetration
21	EN ISO 374-4: 2019	Protective gloves against chemicals
		and micro-organisms - Part 4:
		Determination of resistance to
		degradation by chemicals
22	EN ISO 374-5: 2016	Protective gloves against
		dangerous chemicals and micro-
		organisms - Part 5: Terminology
		and performance requirements for
		micro-organisms risks
23	EN 16523-1: 2015+A1:	Determination of material
	2018	resistance to permeation by
		chemicals - Part 1: Permeationby
		liquid chemical under conditions of
		continuous contact
24	EN ISO 21420: 2020	Protective gloves - General
		requirements and test methods

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.
- Is following the EU-Type Examination 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 the surveillance of the notified body number 2777 by SATRA Technology Europe Ltd.
- This EU Declaration of Conformity is prepared in accordance with Annex IV of Medical Device Regulation (EU) 2017/745.

Authorized Signatory:

Approver : MURAT YILDIZ

Title : General Manager/CEO

MEDIK QU'ANDRIM SIRKETI
Omerli Mah. Separa / Sükrü Koreilu Cad

No:33 - Ali vu / Sylistan Bul
Büyük o Arrace v.D. 626 040 4605

el:0212/48/20 64 Fax: 02/12 / 438 20 65

Approval Date : 15 Feb 2024

Place of Approval : Istanbul, Turkey

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