

EU DECLARATION OF CONFORMITY

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| DOC No. | DOC-MYMEDİKAL-ITC-002 |
| EC Certificate | Not applicable (Self- declared) |
| Manufacturer | MY TICARET VE MEDİKAL A.S. |
| Manufacturer Address | Ömerli mah General Şükrü Koraltı Cd no:33, 34555 Arnavutkoy/Istanbul, Turkey |
| Single Registration Number (SRN) | TR-MF-000018372 |
| Brand | Mumu Guard |
| Product Description | Nitrile Powder Free Examination and Protective 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 | 868302002NPVQ |
| Country of Origin | China |
| Size | XS, S, M, L, XL |
| EAN Code | Blue: 8684266525679, 8684266525686, 8684266525693, 8684266525709, 8684266525716; 8683020022409, 8683020022416, 8683020022423, 8683020022430, 8683020022447 Black: 8683020022669, 8683020022676, 8683020022683, 8683020022690, 8683020022706 |
| European Medical Device Nomenclature (EMDN) | T01020204 (Examination / Treatment Gloves, Nitrile) |
| Global Medical Device Nomenclature (GMDN) | 56286 (Nitrile Examination/Treatment glove, non-powdered, non-sterile) |
| Product Catalogue/ Reference Number | Blue: MGN01-XS, MGN02-S, MGN03-M, MGN04-L, MGN05-XL Black: MGBN01-XS, MGBN02-S, MGBN03-M, MGBN04-L, MGBN05-XL |
| Product Group Reference Number | Blue: SNBE10013-XS, SNBE10014-S, SNBE10015-M, SNBE10016-L, SNBE10017-XL; SNBE20013-XS, SNBE20014-S, SNBE20015-M, SNBE20016-L, SNBE20017-XL |



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| | Black: SNBE20043-XS, SNBE20044-S, SNBE20045-M, SNBE20046-L SNBE20047-XL | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
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| Conformity Assessment Route (MDR): | Annex II and Annex III according to EU 2017/745 | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| Classification & Rule (MDR) | Class I, Rule 1 & Rule 5 according to Annex VIII | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| Device Classification (PPER) | Category III | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| EU Type-Examination Certificate (PPER) | 2777/14815-03/E63-02 | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| Notified Body (PPER) | EU-Type Examination and Ongoing Conformity by Notified Body SATRA TECHNOLOGY EUROPE LTD Bracetown Business Park, Clonee, D15YN2P, Ireland [2777] | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| Applicable Standards | <table border="1"><thead><tr><th>No.</th><th>Regulation/ Standard Number</th><th>Regulation/ Standard Name</th></tr></thead><tbody><tr><td>1</td><td>MDR (EU) 2017/745</td><td>Medical Device Regulation</td></tr><tr><td>2</td><td>PPE (EU) 2016/425</td><td>Personal Protective Equipment Regulation</td></tr><tr><td>3</td><td>ISO 13485: 2016</td><td>Medical devices - Quality management systems - Requirements for regulatory purposes</td></tr><tr><td>4</td><td>ISO 9001: 2015</td><td>Quality management systems – requirements</td></tr><tr><td>5</td><td>ISO 14971: 2019</td><td>Medical devices - application of risk management to medical devices</td></tr><tr><td>6</td><td>EN 455-1: 2020</td><td>Requirements and testing for freedom from holes</td></tr><tr><td>7</td><td>EN 455-2: 2015</td><td>Requirements and testing for physical properties</td></tr><tr><td>8</td><td>EN 455-3: 2015</td><td>Requirements and testing for biological evaluation</td></tr><tr><td>9</td><td>EN 455-4: 2009</td><td>Requirements and testing for shelf-life determination</td></tr><tr><td>10</td><td>ISO 10993-1: 2018</td><td>Biological evaluation of medical devices –Part 1: Evaluation and testing within a risk management process</td></tr><tr><td>11</td><td>ISO 10993-10: 2010</td><td>Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization</td></tr></tbody></table> | No. | Regulation/ Standard Number | Regulation/ Standard Name | 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 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 10993-10: 2010 | Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization |
| No. | Regulation/ Standard Number | Regulation/ Standard Name | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| 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 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 | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
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| 11 | ISO 10993-10: 2010 | Biological evaluation of medical devices — Part 10: Tests for irritation and skin sensitization | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |



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| 12 | ISO 10993-11: 2017 | Biological evaluation of medical devices — Part 11: Tests for systemic toxicity |
| 13 | ISO 20417:2021 | Medical devices - Information to be supplied by the manufacturer |
| 14 | ISO 15223-1: 2021 | Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements |
| 15 | EN ISO 374-1: 2016+A1: 2018 | Protective gloves against dangerous chemicals and micro-organisms - Part 1: Terminology and performance requirements for chemical risks |
| 16 | EN ISO 374-2: 2019 | Protective gloves against dangerous chemicals and micro-organisms - Part 2: Determination of resistance to penetration |
| 17 | EN ISO 374-4: 2019 | Protective gloves against chemicals and micro-organisms - Part 4: Determination of resistance to degradation by chemicals |
| 18 | EN ISO 374-5: 2016 | Protective gloves against dangerous chemicals and micro-organisms - Part 5: Terminology and performance requirements for micro-organisms risks |
| 19 | EN 16523-1: 2015+A1: 2018 | Determination of material resistance to permeation by chemicals - Part 1: Permeation by liquid chemical under conditions of continuous contact |
| 20 | EN ISO 21420:2020 | Protective gloves – General requirements and test methods |
| 21 | ASTM D 6978-05:2019 | Standard Practice for Assessment of Resistance of Medical Gloves to Permeation by Chemotherapy Drugs |
| 22 | ASTMF1671/F1671-13 | 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 System |



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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 documentation is retained under the premise of the manufacturer.
- The gloves are manufactured according to EN ISO 9001:2015 and EN ISO 13485:2016 Quality Management System.
- Is following to 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 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 TICARET VE
MEDİKAL ANONİM ŞİRKETİ
: Ömerli Mah. General Şükrü Koraltı Cad
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Approval Date : 02 Sep 2024

Place of Approval : Istanbul, Turkey

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