

DECLARATION OF CONFORMITY

Manufacturer : St Marys Rubbers Pvt. Ltd
XVII /401A, Thottamkavala, Vizhikkathode, Koovappally PO
Kanjirappally, Kottayam, Kerala- 686518, India

SRN : IN-MF-000008817

European Representative : Emergo Europe, Westervoortsedijk 60, 6827 AT Arnhem ,The Netherlands
SRN : NL-AR-000000116

Device Name : Sterile Latex Surgical Gloves Powdered
Brand : Medismart, Medistar, Mediten, One Glove, Medismart Premium, Mediagni
Size : 5.5, 6.0, 6.5, 7.0, 7.5, 8.0, 8.5 & 9.0

Device Intended use : Sterile Latex Surgical Gloves intended for single use on one individual during single surgical procedure and to be worn once and then discarded. The gloves provide a protective barrier that reduces the risk of contamination and transmission of infectious agents, microorganisms, such as bacteria and viruses during medical procedures , particularly surgical interventions.

The gloves are designed to protect the patient from cross-contamination and to minimize the risk of post-operative infections. The glove is pre powdered with absorbable corn starch for easy donning.

Basic UDI-DI : 8908004933SLSP1MZ
EMDN Code : T01010101 – Surgical Gloves, Latex, Powdered
MDA/MDN Code : MDN 1204
MDS & MDT code(s) : MDS 1005, MDT 2011, MDT 2006, MDT 2008
Sterilization Method : ETO sterilization/ Gamma Sterilization
Batch No :
Product reference number :

1. Medical Device details

Device Classification : Class IIa, Rule 06 as per MDCG 2021-24 & Annex VIII of MDR 2017/745
Conformity Assessment Route : Annex IX

Declaration : We hereby under our sole responsibility declare that the above mentioned product meets the provisions of the Medical Device Regulation EU 2017/745, applicable harmonized standards, and as transposed into national laws. All supporting documentation is retained under the premises of the manufacturer.



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Standards Applied

: Applicable Harmonized Standards

EN ISO 13485:2016/A11:2021, EN ISO 14971:2019/A11:2021, EN ISO 20417:2021, EN ISO 15223-1:2021, EN 62366-1:2015+AC:2015+AC:2016+A1:2020, EN ISO - 10993-1:2020, EN ISO 10993-5:2009, EN ISO 10993-7:2008/AC:2009, EN ISO 10993-10:2023, EN ISO 10993-11:201, EN ISO 10993-12:2021, EN ISO 10993-23:2021, EN ISO 11135:2014/A1:2019, EN ISO 11138-2:2017, EN ISO 11737-1:2018/A1:2021, EN ISO 11737-2:2020, EN ISO 11607-1:2020, EN ISO 11607-2:2020+A1:2023, EN 455-1:2020+A2: 2024, EN 455-2:2024, EN 455-3:2023, EN 455-4:2009, EN 556-1:2024

: Applicable Other Standards

ISO 13485:2016, ISO 14971:2019, ISO 15223-1:2021, IEC 62366-1:2015/AMD1:2020 +COR1:2016, ISO 10993-1:2018, ISO 10993-5:2009, ISO 10993-7:2008/AMD1: 2019 +COR 1:2009, ISO 10993-10:2021, ISO 10993-11:2017, ISO 10993-12:2021, ISO 10993-23:2021, ISO 11135:2014/AMD 1: 2018, ISO 11138-2:2017, ISO 11737-1:2018/AMD1: 2021, ISO 11737-2:2019, ISO 11607-1:2019, ISO 11607-2:2019, ISO 10282:2023, ASTM D 3577-19(2023), ASTM F1980-21, ASTM D 6124-06:2022, ASTM D5712-15:2020, ASTM D4169:2016, ASTM D999:2008, ISO 2233:2000, ISO 2248:1985, ISO 2247:2000, IS 7028-1:2002, IS 13422:2024(ISO 10282:2023), ISO 14644-1:2015, ISO 14644-2:2015, ISO

14644-4:2022

: Guidelines

MEDDEV 2.5/9 Rev 1, MEDDEV 2.7.1 Rev 4, MEDDEV 2.4/1 Rev 9, MEDDEV 2.12/2 Rev 2, MDR 2017/745, EU 2024/1860

Notified Body

: DNV Product Assurance AS
Veritasveien 1, 1363 Høvik, Norway

Notified Body No

: 2460

EC Certificate No.

: C556189

Date & place of Issue

: 06 February 2025, Høvik

Validity date

: 05 February 2030

2. Personal Protective equipment Details

Device Classification

: Personal Protective Equipment Category III as per regulation (EU) 2016/425

Conformity Assessment

Route

: EU type Examination (Module B) set out in Annexure V and conformity to type based on quality assurance of the production process (Module D) set out in Annexure VIII.

PPE Standards

: EN ISO 374-1:2016 + A1:2018(type B), EN ISO 374-5:2016, EN ISO 374-2:2019, EN ISO 374-4:2019, EN 16523-1:2015+A1:2018, ISO 21420:2020, ISO 3071:2020, ISO

16604:2004



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Applicable Guidance Documents : EU 2016/425

Declaration : We declare under our sole responsibility that the above mentioned product complies with the essential requirements of PPE Regulation (EU) 2016/425. All Prior amendments are and as transposed into national laws.

Name of Notified Body& Address : SGS Fimko Oy
: Notiifed Body 0598, Takomotie 8, FI-00380 Helsinki, Finland
Notified Body Number : 0598

I. EU Type –Examination Certificate, Module B

Certificate No. : 0598/PPE/24/5509 Issue 1
Date & Place of Issue : 06 December 2024, SGS FIMKO Ltd
Validity Date : 06 December 2029

II. Module D certificate details

Certificate No. : IN22/00000837
Date & Place of Issue : 15 December 2022, SGS FIMKO Ltd
Validity Date : 15 December 2025

Place & date of Issue : Kanjirappally, 22.09.2025

Signature

Name & Designation



: Rony Emmanuel
Manager QA & PRRC

