Innovation & Quality

EU DECLARATION OF CONFORMITY

Manufacturer : Hartalega NGC Sdn. Bhd.

Manufacturer's Address : No.1, Persiaran Tanjung, Kawasan Perindustrian Tanjung,

43900 Sepang, Selangor Darul Ehsan, Malaysia.

EU Representative : MDSS GmbH

Schiffgraben 41, 30175 Hannover, Germany. Reference Only

Product Description (EU MDR) : Nitrile Powder Free Examination Glove

Device Classification (EU MDR) : Class I, according to Annex VIII of Regulation (EU) 2017/745

Rule(s) : 1 and 5

Conformity Assessment : Annex II and Annex III

Procedure

Basic UDI-DI : 955100777HNGCTFMD005AQ8

Authorised Representative SRN : DE-AR-000005430

Manufacturer SRN : MY-MF-000010459

Product Description (EU PPER) : _ Nitrile Powder Free Examination Gloves (≥ 2.2 mil)

Available in standard minimum 240mm length or a longer

cuff variant of 280mm

Device Classification (EU PPER) : Category III (Type B)

EU Type-Examination : 2777/11578-03/E00-00

For Roll Intended Purpose For Rolling: Nitrile Powder Free Examination Glove are intended to be

used to contribute to prevent cross contaminations in the framework of medical examinations and diagnostic/therapeutic procedures conducted under non-sterile

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conditions.

and

Nitrile Powder Free Examination Glove is intended to protect users from substances and mixtures which are hazardous to

health and harmful biological agents that may cause very

serious consequences or irreversible damage to health.

Standard Reference : Attachment I

Reference to Trade Name Parent: Attachment II Reference Only

Hartalega Holdings Berhad (741883-X)

Certificate Number (EU PPER)

C-G-9, Jalan Dataran SD1, Dataran SD PJU 9 Bandar Sri Damansara 52200 Kuala Lumpur, Malaysia Tel: +603 - 6277 1733 Fax: +603 - 6280 2533 www.hartalega.com.my Hartalega NGC Sdn Bhd (984586-P)

No. 1, Persiaran Tanjung Kawasan Perindustrian Tanjung 43900 Sepang, Selangor, Malaysia Tel: +603 - 8707 3000 Rev 24

Growing Global

We, Hartalega NGC Sdn. Bhd. herewith declared that above mentioned device: • is in conformity with the Regulation (EU) 2017/745 of The European Parliament and of The Council of medical devices. is in conformity with the provisions of Regulation (EU) 2016/425 on personal protective For Reference equipment. is subject to the conformity assessment procedure Module D set out in Annex VIII of Regulation (EU) 2016/425, under the surveillance of the notified body SATRA Technology Europe Limited, Bracetown Business Park, Clonee D15YN2P, Republic of Ireland (Notified eterence Only Body number 2777). This EU declaration of conformity is issued under the sole responsibility of the manufacturer, Hartalega NGC Sdn. Bhd. : Hartalega NGC Sdn. Bhd./ 24th May 2024 Place and Date of Issue For Reference Only For Refer Signed on Behalf of Hartalega NGC: For Reference Only For Reference Only For Reference Only For Reference Only Name or: NURUL ALSYAH KONG Position: GENERAL MANAGER - QUALITY ASSURANCE nce Only For Reference Only For Reference For Reference Only For Reference Only For Reference Only

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| Standard | non Only For Reference On |
|---|---|
| ISO 9001:2015 | Quality Management Systems – Requirements |
| EN ISO 13485:2016+A11:2021 | Medical Devices – Quality Management Systems – Requirements for Regulatory Purposes |
| EN 455-1:2020+A1:2022 | Medical Gloves for Single Use Part 1: Requirements and Testing for Freedom from Holes |
| BS EN 455-1:2020+A1:2022 | Medical Gloves for Single Use Part 1: Requirements and Testing for Freedom from Holes |
| EN 455-2:2015 | Medical Gloves for Single Use Part 2: Requirements and Testing for Physical Properties |
| BS EN 455-2:2015 | Medical Gloves for Single Use Part 2: Requirements and Testing for Physical Properties |
| EN 455-3:2023 | Medical Gloves for Single Use Part 3: Requirements and Testing for Biological Evaluation |
| BS EN 455-3:2023 | Medical Gloves for Single Use Part 3: Requirements and Testing for Biological Evaluation |
| EN 455-4:2009 nce Only | Medical Gloves for Single Use Part 4: Requirements and Testing for Shelf Life noe Only Determination |
| nce Only For Refere BS EN 455-4:2009 For Reference Only | Medical Gloves for Single Use — Control For Reference Or Part 4: Requirements and Testing for Shelf Life Determination — Control For Reference Only |
| BS EN ISO 20417:2021 | Medical devices – Information to be Supplied by the Manufacturer |
| BS EN ISO 14971:2019+A11:2021 | Medical Devices – Application of Risk Management to Medical Devices – Application of Risk Management to |
| ISO 15223-1:2021 or Refere | Medical Devices –Symbol to be Used with Information to be Supplied by the Manufacturer — Control Part 1: General Requirements |
| For Reference Only ISO 10993-1:2018 | Biological Evaluation of Medical Devices — Errence Control For Res Part 1: Evaluation and Testing within a Risk Management Process — Errence Or |
| BS EN ISO 10993 - 1: 2020 | Biological Evaluation of Medical Devices Part 1: Evaluation and Testing within a Risk Management Process |
| ISO 10993-5:2009 | Biological Evaluation of Medical Devices Approximately For Reference Or Part 5: Tests for In Vitro Cytotoxicity |

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| Standard | Title | |
|--------------------------------------|---|--|
| BS EN ISO 10993 – 5:2009 | Biological Evaluation of Medical Devices of a Comby Part 5: Tests for In Vitro Cytotoxicity | |
| ISO 10993-10:2021 | Biological Evaluation of Medical Devices Part 10: Tests for Skin Sensitization | |
| BS EN ISO 10993 – 10:2023 | Biological Evaluation of Medical Devices Part 10: Tests for Skin Sensitization | |
| ISO 10993-11:2017 | Biological Evaluation of Medical Devices Part 11: Tests for Systemic Toxicity | |
| BS EN ISO 10993 – 11:2018 | Biological Evaluation of Medical Devices Part 11: Tests for Systemic Toxicity | |
| ISO 10993-18:2020/ Amd 1:2022 | Biological Evaluation of Medical Devices Part 18: Chemical Characterization of Medical Device Materials Within A Risk Management Process Amendment 1: Determination of the uncertainty factor | |
| BS EN ISO 10993 – 18:2020+A1:2023 | Part 18: Chemical Characterization of Medical Device | |
| ISO 10993-23:2021 | Biological Evaluation of Medical Devices Part 23: Tests for Irritation | |
| BS EN ISO 10993 – 23:2021 | Biological Evaluation of Medical Devices Part 23: Tests for Irritation | |
| ISO 2859-1:1999/Amd1:2011 | Limit (AQL) for Lot-By-Lot Inspection | |
| ASTM D4169-22 | Standard Practice for Performance Testing of Shipping Containers and Systems | |

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| Standard | For Reference Only Title or Reference Only | For Refer |
|---------------------------|---|------------------------|
| EN ISO 21420:2020 | Protective gloves - General requirements and test methods | ence Only |
| EN ISO 374-1:2016+A1:2018 | Protective gloves against dangerous chemicals and micro- organisms - Part 1: Terminology and performance requirements for chemical risks | For Refer |
| EN ISO 374-5:2016 | Protective gloves against dangerous chemicals and micro- organisms - Part 5: Terminology and performance requirements for micro- organisms risks | ence Only For Refer |
| EN 421:2010 | Protective gloves against ionizing radiation and radioactive contamination | ence Only |

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ATTACHMENT II

| | For Reference Only For Reference Only For Reference Only | | For Refer |
|------------|--|--|-----------|
| | Product or Trade Name | Reference Number | |
| | ence Only For Reference Only | For Referer XS: 436XS For Refer S: 436S | |
| | PEPPLER NITRIL CLOUD 436 | ence Only L: 436L XL: 436XL | |
| | For Reperler nitril black 444 Refer | XS: 444XS S: 444S ence Only L: 444L | For Refer |
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