

CE EU 2017/745 Medical Device Regulation



Declaration of Conformity – EU

Maxter Glove Manufacturing Sdn Bhd hereby confirms that the product mentioned below complies with EU Regulations and Standards and is manufactured according to ISO9001 & ISO13485 standard requirements.

Aurelia Absolute 100 3.2 mil Black Powder Free Nitrile Examination Glove

Size	Product Code	Barcode
X-Small	9899A5	955-500211-9775
Small	9899A6	955-500211-9782
Medium	9899A7	955-500211-9799
Large	9899A8	955-500211-9805
X-Large	9899A9	955-500211-9812

Classification of the product:

- Class I Medical Device based on Rule 5 transient use, Annex VIII of Medical Device Regulation (EU) 2017/745
- CAT III PPE (EU) 2016/425

Product mentioned above complies with:

- The General Safety and Performance requirements of Annex I, Medical Regulation (EU)2017/745 for Class I Medical Devices and with the Article 19 requirements
- The provisions of Personal Protective Equipment (PPE) Regulation (EU) 2016/425, including the General Safety Requirements (Annex II), Module B EU-Type Examination Certification and Module D, Conformity to type, based on Quality Assurance of the production process.
- EEC regulations concerning the conformity of materials and products that are allowed to come into contact with food. In accordance with Regulation EEC 1935/2004, Regulation EC 10/2011 & Regulation EC 2023/2006.
- Medical Device Regulation 2002 for Class I Medical Devices

Certification:

- Module B, EU Type Examination Certificate issued by Notified Body: Satra (2777) Certificate No. 2777/12716-02/E00-00
- Module D, Regulation EU 2016/425, Examination Certificate issued by Notified Body: SGS FIMKO OY, Notified Body CE0598 – Certificate No. MY19/1811030073
- ISO9001:2015
- ISO13485:2016

Gloves tested according to Harmonised Standards:

- EN374-1 chemical resistance
- EN374-5 microbiological resistance
- EN455 1,2,3, 4 medical devices
- EN21420- physical attributes or EN420 during the transition period



User Information:

- The gloves are suitable for contact with dry, fatty, alcoholic, and aqueous food for short term contact based on the outcome of the overall migration test on the food simulants.
- The product does not contain natural rubber latex. Contains accelerators which may cause allergic reactions. Please retain the packaging for reference.
- Store below 40°C/104°F in dry, clean condition and away from direct sunlight.

Responsibility

 This Declaration of Conformity is issued under the responsibility of the Manufacturer as indicated below:

Manufacturer:

- Maxter Glove Manufacturing Sdn Bhd, located at Lot 6070, Jalan Haji Abdul Manam, 6th Miles off Jalan Meru, 41050 Klang, Selangor, Malaysia
- SRN: MY-MF-000016719

Importer & Authorised Representatives:

- EU Representative is Supermax Healthcare (Europe) Limited, 38 Main Street, Swords Co. Dublin, K67 E0A2, Ireland
 - ♣ SRN: IE-AR-000013888
- UK Representative is Supermax Healthcare Limited, 12-16 Titan Drive, Fengate, PE1 5XN, Peterborough, United Kingdom

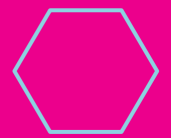
Maxter Glove Manufacturing Sdn Bhd, Malaysia

Authorised by:

Yap Peak Geeh Senior Manager QA & Regulatory Affairs

Date: 27-May-2022

*This declaration is valid for period of 2 years from the date of issue or until any changes to regulations or products are applicable.



UKCA
UK Medical Device
Regulation 2002



Declaration of Conformity – UK

Maxter Glove Manufacturing Sdn Bhd hereby confirms that the product mentioned below complies with UK Regulations and Standards and is manufactured according to ISO9001 & ISO13485 standard requirements.

Aurelia Absolute 100 3.2 mil Black Powder Free Nitrile Examination Glove

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X-Small	9899A5	955-500211-9775
Small	9899A6	955-500211-9782
Medium	9899A7	955-500211-9799
Large	9899A8	955-500211-9805
X-Large	9899A9	955-500211-9812

Classification of the product:

- Class I Medical Device based on Rule 5 transient use, Annex IX of COUNCIL DIRECTIVE 93/42/EEC
 of 14 June 1993 concerning medical devices, which is implemented into UK law by the Medical
 Device Regulations 2002, as amended.
- Class I Medical Device based on Rule 5 transient use, Annex VIII of Medical Device Regulation (EU) 2017/745
- CAT III PPE (EU) 2016/425, which is directly implemented into UK law.

Product mentioned above complies with:

- Medical Device Regulation 2002 for Class I Medical Devices, implementing COUNCIL DIRECTIVE 93/42/EEC
- The General Safety and Performance requirements of Annex I, Medical Regulation (EU) 2017/745 for Class I Medical Devices and with the Article 19 requirements.
- The provisions of Personal Protective Equipment (PPE) Regulation (EU) 2016/425, including the General Safety Requirements (Annex II), Module B EU-Type Examination Certification and Module D, Conformity to type, based on Quality Assurance of the production process.
- EEC regulations concerning the conformity of materials and products that are allowed to come into contact with food. In accordance with Regulation EEC 1935/2004, Regulation EC 10/2011 & Regulation EC 2023/2006.

Certification:

- Module B, UKCA Type Examination Certificate issued by Approved Body: Satra (0321) Certificate No. AB0321/19868-02/E00-00
- Module D, Regulation 2016/425 as brought into the UK Law, Examination Certificate issued by Approved Body: SGS UK, Approved Body UKCA0120 – Certificate No. MY21/1811030640
- ISO9001:2015
- ISO13485:2016



Gloves tested according to Harmonised Standards:

- EN374-1 chemical resistance
- EN374-5 microbiological resistance
- EN455 1,2,3, 4 medical devices
- EN21420- physical attributes or EN420 during the transition period

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- EU Representative is Supermax Healthcare Europe Limited, 38 Main Street, Swords Co. Dublin, K67
 E0A2, Ireland
- UK Representative is Supermax Healthcare Limited, 12-16 Titan Drive, Fengate, PE1 5XN, Peterborough, United Kingdom

Maxter Glove Manufacturing Sdn Bhd, Malaysia,

Authorised by:

Yap Peak Geeh Senior Manager

QA & Regulatory Affairs

Date: 25-Jun-2022

*This declaration is valid for period of 2 years from the date of issue or until any changes to regulations or products are applicable.