



Personal Protective Equipment Regulation (EU) 2016/425

Quality Assurance Certificate

Module D

Manufacturer

Unimax Medical Products Co., Ltd.

Special No.1 Liansai Road, Changshangkou Town, 433024, Xiantao City, Hubei Province
People's Republic of China

Scope:

Chemical Protective Clothing – Single Use Coveralls

Certificate Number: 54185

Issued by: BTTG™ (Notified Body No. 0338 for Regulation (EU) 2016/425)

First issue: 11 October 2019 **Date of Issue:** 22 October 2019 **Expiry*:** 11 October 2022

Authorised by

C A Butcher
Certification Manager

*Subject to continued compliance and audit.

The attached schedule of approval forms part of this certificate.

Note: The validity of this certificate can be confirmed by contacting the Issuing Office:
BTTG™, Unit 6 Wheel Forge Way, Trafford Park, Manchester, M17 1EH, United Kingdom
Tel: +44 (0)161 876 4211 email: ppe@bttg.co.uk website: www.bttg.co.uk



Schedule of Approval Certificate Number: 54185

First Issued:	11 October 2019	Page No:	2 of 3
Issue date:	22 October 2019	Issued by:	BTTG™ (Notified Body No. 0338)
Expiry date:	11 October 2022	BTTG™ ref:	E-006644

Manufacturer: Unimax Medical Products Co., Ltd.

BTTG™, specified as a "notified body" under the terms of the Regulation (EU) 2016/425 of the European Parliament and of the Council on personal protective equipment, did undertake the relevant quality assessment procedures for the manufacturing sites identified below, which were found to be in compliance with Module D "conformity to type based on quality assurance of the production process" of the Regulation, subject to any conditions in the schedule attached hereto.

This certificate authorises the use of the Mark of Conformity (the 'CE mark'), together with the number of the Notified Body involved in the production control phase (0338) once the manufacturer has issued a Declaration of Conformity according to Article 15 of the Regulation.

Places of Production

Special No.1 Liansai Road, Changshangkou Town, 433024, Xintao City, Hubei Province
People's Republic of China

Approval Documents

BTTG/JY/Unimax/04, 2018

CONTINUED ON PAGE 3



Schedule of Approval Certificate Number: 54185

First Issued:	11 October 2019	Page No:	3 of 3
Issue date:	22 October 2019	Issued by:	BTTG™ (Notified Body No. 0338)
Expiry date:	11 October 2022	BTTG™ ref:	E-006644

Manufacturer: Unimax Medical Products Co., Ltd.

Terms and Conditions associated with the issue of this Quality Assurance Certificate

1. This certificate is issued subject to BTTG™'s standard terms of business.
2. Production is limited to the site(s) listed above.
3. The client must implement appropriate changes as notified by BTTG™.
4. The client must ensure the certified product is representative of the ongoing manufactured product.
5. The client must:
 - a) Permit ongoing surveillance and access to documentation and records, and access to the relevant equipment, location(s), area(s), personnel and clients subcontractors.
 - b) Investigate complaints associated with the certified products. Records of such complaints, and actions taken, must be kept by the client and made available to BTTG™ when requested.
 - c) Allow participation of observers during surveillance audits when requested.
6. The client must only make claims consistent with the scope of certification,
7. The client must not make any misleading or unauthorised comments regarding the certified product or the certification body.
8. The client must upon suspension, withdrawal, or termination of certification discontinue the use of all advertising matter that contains any reference thereto and take action to return this certificate to BTTG™.
9. The client must comply with the requirements for the use of the notified body number as detailed below.
10. Changes to a client's product design, manufacturing processes, operations, location, management team or resource provision that could have an impact on the certified product shall be immediately notified to BTTG™.
11. This certificate is issued in the English language only. It is the responsibility of the Manufacturer / Authorised Representative to obtain and supply language versions acceptable to the country where the product is to be sold.
12. This certificate remains the property of BTTG™ and will be withdrawn if any of the conditions attached to its issue are not complied with.
13. The Manufacturer shall have continuous surveillance of Factory Production Control carried out by a Notified Body and a re-certification of Factory Production Control every three years.

Use of Notified Body Number

1. The Notified Body Number must only be used:
 - a) In direct association with products or systems covered by this Quality System Certificate.
 - b) By holder(s) of the Certificate.
2. Use of BTTG™ Notified Body Number does not extend to other companies which are:
 - a) part of the same corporate group as the Certificate holding company: or
 - b) named in a Certificate, for example as a supplier.
3. Particular care must always be taken to avoid the association of the BTTG™ Notified Body Number with other products or systems or schemes and with claims or information not contained in the BTTG™ document.
4. The EC mark consists of the letters 'CE', in the form given in Annex II of Regulation (EC) No. 765/2008, followed by BTTG™'s notified body number.

If any of the above requirements are not met BTTG™ will seek to suspend, withdraw or terminate this certificate.

END OF REPORT