

# **CERTIFICATE LETTER**

This is to certify that INSPEC International B.V., Notified Body 2849, has accepted transfer of type-examination (Module B) and/or approval decision (Module C2 or D) certification from INSPEC International Ltd, Notified Body 0194 in accordance with the transitional arrangements of the withdrawal of the United Kingdom of Great Britain and Northern Ireland from the European Union and that the original Certificate(s) combined with this Certificate Letter evidences that INSPEC International B.V. deems them to be in compliance with the Personal Protective Equipment Regulation (EU) 2016/425.

Manufacturer: Medop S.A.

Zabala 16-4º izq 48003 Bilbao Vizcaya Spain

The scope of the certification is for the Transferred Certificate(s) listed on page 2.

Where a type-examination certificate has been transferred, this certificate, while valid, serves as authorisation to the manufacturer to reference INSPEC International B.V. within information supplied to the user, and their EU Declaration of Conformity.

Where an Approval Decision (Module C2 or D) has been transferred, this certificate, while valid, serves as authorisation to the manufacturer to affix our notified body identification number, 2849, to each individual item of PPE that is in conformity with the type described in the type-examination certificate(s) for all production beyond the Date of current issue.

The Date of expiry of certification for the Type and / or Approval Decision is stated within the Transferred Certificate(s) listed on page 2. Where no expiry date is shown, the certificate shall expire on 21 April 2023 as per the transitional provisions of Article 47(2).

Date of initial certification: 30 November 2020 Date of current issue: 30 November 2020

Certification Manager







# **Transferred Certificates**

The following is a list of original certification documentation issued by INSPEC International Ltd, Notified Body 0194 which were transferred without revision to INSPEC International B.V., Notified Body 2849 and are confirmed as valid when combined with this certificate letter.

1198	PPE19161288	PPE19161380
1660	PPE19161289	PPE19161393
3064	PPE19161290	PPE19161705
969	PPE19161352	PPE20161895
D-PPE20171083	PPE19161353	PPE20161896
PPE18161103	PPE19161354	PPE20161897
PPE19161285	PPE19161364	PPE20161919
PPE19161286	PPE19161378	PPE20161955
PPE19161287	PPE19161379	PPE20161957

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### Certificate amendment record

Date	Description
30/11/2020	Initial Issue

#### General Conditions attached to the issue of this certificate:

- The manufacturer / authorised representative shall undertake to fulfil the obligations arising out
  of the Personal Protective Equipment Regulation (EU) 2016/425, and with INSPEC's
  Regulations governing the Module as displayed on original Certificates.
- 2. Personal Protective Equipment Directive, 89/686/EEC, Article 10 certificates and Article 11.B approval decisions are valid under Personal Protective Equipment Regulation (EU) 2016/425, Article 47(2).
- 3. INSPEC International Ltd, Notified Body 0194, shall be absorbed into INSPEC International B.V. and closed this is to be reflected on the EU NANDO website as "(ex-0194)" following the Body type "NB 2849". Therefore INSPEC International B.V. shall be available to verify certificates and references to the closed Notified Body 0194.
- 4. INSPEC International Ltd have requested that manufactures cease making reference to INSPEC International Ltd, Notified Body 0194 as soon as is practical, and to a date not later than 30 May 2021 for any production on or beyond 01 January 2021.
- 5. This certificate remains the property of INSPEC and may be withdrawn if any of the conditions attached to its issue are not complied with.
- 6. This certificate may be copied or reproduced by the certificate holder, complete and accompanied by the applicable Transferred Certificate(s), and without omissions or additions. Their use must be in accordance with INSPEC's terms of business.

## Module B: Conditions attached to the issue of this certificate:

- 7. The manufacturer / authorised representative shall inform INSPEC of any planned changes to the product or technical file which may affect the validity of this certificate, before any such change is made.
- 8. Marking and instructions have been assessed in the English language only. It is the manufacturer's / authorised representative's responsibility to obtain and supply language versions acceptable to the country where the product is to be sold.
- 9. For category III product, the manufacturer must obtain and maintain an approval decision to Module C2 or Module D prior to placing product on the Union market.

INSPEC International B.V. • Beechavenue 54-62 • 1119 PW • Schiphol-Rijk • The Netherlands.• Notified Body 2849

This certificate has been issued in accordance with our standard terms and conditions and is subject to INSPEC Regulations and Conditions of Use. This certificate remains the property of INSPEC and may be withdrawn if any of the conditions attached to its issue are not complied with by the certified company.

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# Module C2: Conditions attached to the issue of this certificate

- 10. The manufacturer / authorised representative shall inform INSPEC of any planned changes to the product type which may affect the validity of this certificate.
- 11. For the certificate to remain valid a minimum of annual sampling to perform product checks must be conducted, as per Annex VII, 4.
- 12. The manufacturer may affix INSPEC's notified body identification number, 2849, to each PPE, and draw up a written EU declaration of conformity for each PPE model referencing this certificate and the Type-examination certificate(s) as per Annex VII, 6.

### Module D: Conditions attached to the issue of this certificate

- 13. The manufacturer / authorised representative shall undertake to fulfil the obligations arising out of the quality system, and of the Personal Protective Equipment Regulation (EU) 2016/425, Annex VIII, and with INSPEC's Regulations governing this Module.
- 14. The manufacturer / authorised representative shall inform INSPEC of any planned changes to the quality system, wherein INSPEC will proceed with evaluation of the proposals as per Annex VIII, 3.5.
- 15. For the certificate to remain valid audits and visits must be conducted, as per Annex VIII, 4.
- 16. The manufacturer may affix INSPEC's notified body identification number, 2849, to each PPE, and draw up a written EU declaration of conformity for each PPE model referencing this certificate and the Type-examination certificate(s) as per Annex VIII, 5.

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