

# Conformity to Type based on Quality Assurance of the Production Process

This is to certify that:

Supreme Visors Limited  
Unit 2  
Aston Fields Road  
Whitehouse Industrial Estate  
Runcorn  
WA7 3DL  
United Kingdom

Holds Certificate Number:

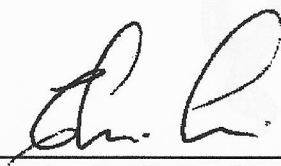
CE 575934

In respect of:

**The manufacture of Respiratory Protective Equipment to the standards specified on the continuation sheet.**

on the basis BSI carried out the quality assurance assessment in accordance with the Regulation (EU) 2016/425 of the European Parliament and Council relating to Personal Protective Equipment Regulation (PPE) Annex VIII

For and on behalf of BSI, a Notified Body for the above Regulation (Notified Body Number 0086):



Chris Lewis - Certification Director, Product Certification

First Issued: 2011-09-14

Latest Issue: 2019-02-11

Effective Date: 2019-02-11

Expiry Date: 2024-02-11

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...making excellence a habit.™

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To check its validity telephone +44 (0) 345 080 9000. An electronic certificate can be authenticated [online](#).

BSI Assurance UK Limited, registered in England under number 7805321 at 389 Chiswick High Road, London W4 4AL, UK.  
A member of BSI Group of Companies.

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No. CE 575934

## Product Specifications

The Respiratory Protective Equipment covered by the Module D scope of this Certificate conform to the following standards:

### Standard

EN 14594:2005

### Product Type

Respiratory protective devices. Continuous flow compressed air line breathing apparatus

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**Certificate Administration Details.**

**Certificate Amendment Record**

Issue date	Comments	BSI Project Number
February 2019	Transition of certificate to Module D under PPE Regulation (EU) 2016/425, formerly Article 11b of the PPE Directive 89/686/EEC.	0086:19:9717077

**Certificate validity**

The Certificate holder is responsible for ensuring that the Notified Body is advised of changes to any aspect of the overall process utilised in the manufacture of the products, failure to do so could invalidate the Certificate in respect of product manufactured following the introduction of such changes

The validity of the Certificate is also dependent on the maintenance of an ISO 9001 quality system certified by a recognized certification organisation

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