



Declaration of Conformity

Language of this declaration: **en** English (translations available in the following pages)

EU Declaration of conformity n°
Revision n°
Technical file n°

DC Prevention Plus Boot Cover
01
02 A - 26

| | |
|---------------------------|--|
| Legal manufacturer | Medline International France SAS 5 rue Charles Lindbergh 44110 Châteaubriant - France |
| EU representative | <p>Single Registration Number Not available</p> <p>Not applicable</p> <p>Single Registration Number Not applicable</p> |
| Product type | Protective clothing - Prevention Plus |
| Product Code(s) | NONE27348PXL |
| GMDN Code(s) | 13576 |

European Union Regulations:

We herewith declare under our sole responsibility that the above-mentioned products meet the provisions of the following EU Regulations and/or Council Directive(s) as transposed into national laws.

| | |
|--|---|
| Applicable directive : | Medical Device: Directive 93/42/EEC of 14 June 1993 as amended. |
| Medical Device classification | Class I; Rule n°1 |
| Applicable standards and/or Common Specifications are listed in technical file n°: | 02 A |
| Conformity assessment procedure Certificate n° Notified Body (name/number) | Annex VII Not applicable, self Declaration Not applicable, self Declaration |

| | |
|--|--|
| Applicable regulation: | Personal Protective Equipment: Regulation (EU) 2016/425 of 9 March 2016 |
| Risk categories of PPE | Category III; Risk: Substances and mixtures which are hazardous to health |
| Applicable standards are listed in technical file n° | 26 |
| Conformity assessment procedure(s) | |
| <input type="checkbox"/> Module A set out in Annex IV of Regulation <input checked="" type="checkbox"/> Module B set out in Annex V of Regulation EU-type examination Certificate n° Notified Body (name/number) <input type="checkbox"/> Module C set out in Annex VI of Regulation | CE 706248 BSI (2797) |
| Where applicable, the PPE is subject to the following conformity assessment procedure under surveillance of the notified body (name, number): | BSI (2797) |
| <input checked="" type="checkbox"/> Module C2 set out in Annex VII of Regulation : Conformity to type based on internal production control plus supervised product checks at random intervals <input type="checkbox"/> Module D set out in Annex VIII of Regulation : Conformity to type based on quality assurance of the production process | |

Australian Regulations:

This is a declaration of conformity made under clause 6.6 of Schedule 3 to the Therapeutic Goods (Medical Devices) Regulations 2002.
Each kind of medical device to which the technical documentation applies complies with the applicable provisions of the essential principles and the classification rules before being supplied.

Authorised Signatory:

Kenneth Smith
Senior QA/RA Manager

44110 Châteaubriant - France
Place

Date

FINT.520 Revision n° 05 Effective date: 15 July 19

| | |
|-----------|---|
| 05.INT.02 | Duration of archiving: 10 years after the end of life |
|-----------|---|

See intranet or the server for latest released version. For internal use only. This document contains confidential, proprietary information of Medline Int. or one of its subsidiaries. It may not be copied or reproduced without prior written permission from Medline Int.