



## **Declaration of Conformity for Medline US products**

Declaration of conformity no

Revision no

Technical file #

DC Sensicare PI micro

01

CE2009005 (MDD) and CEP2009005 (PPE)

Legal Manufacturer Medline Industries, Inc.

Three Lake Drive, Northfield, illinois 60093 USA

**EC Representative** Medline International France SAS

5 rue Charles Lindbergh, 44110 Châteaubriant - France

Product range Surgical Gloves - Sensicare PI micro

**Product codes** See attached list

Classification Medical Device class IIa Sterile - rule 6

Personal Protective Equipment Category III

**GMDN Codes** 56293 Polyisoprene surgical glove, non-powdered

#### **European Union Regulations:**

#### **European Representative**

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

Medical Devices Directive: Council Directive 93/42/EEC of 14 June, 1993 as amended. Applicable directive:

Annex 93/42/EEC Annex II

BSI The united Kingdom - CE 0086 **Notified Body** 

Certificate no CE 557201

The United Kingdom, 20th January 2010 First Issued (Place/Date)

See Technical File CE2009005 Applicable standards

Personal Protective Equipment: Council Directive 89/686/EEC of 21 December 1989. Applicable directive:

#### EC-type examination per Article 10

Certificate nº CE 664226

Notified Body (name/number/address) BSI The united Kingdom - CE 0086 The United Kingdom, 21st March 2017 First Issued (Place/Date)

Applicable standards See Technical File CEP2009005

#### Conformity assessment procedure per Article 11

Article 11 A or B 11A

Certificate no CE 651845

Notified Body (name/number/address) BSI The united Kingdom - CE 0086 First Issued (Place/Date) The United Kingdom, 8th june 2016

#### Australian Regulations:

This is a declaration of conformity made under clause 1.8 of Schedule 3 to the Therapeutic Goods (Medical Devices) Regulations 2002.

Each kind of medical device to which the system has been applied complies with the applicable provisions of the essential principles, the classification rules, and the full quality assurance procedures, at each stage, from the design of the device until its final inspection before being supplied.

All supporting documentation is retained at the premises of the manufacturer.

Authorised Signatory:

**Kenneth Smith** 

International Quality Operations Manager

44110 Châteaubriant

Duration of archiving: End of life and support period

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# List of Medline US finished products for declaration of conformity

**FINT.491**Revision n°00
Effective date: 21 Oct 16

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**Surgical gloves - Sensicare PI micro** 

**DoC nr:** DC Sensicar **Rev:** 01

Product number	Description	GMDN Code	Class	PPE Class
MSG9655	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9660	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9665	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9670	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9675	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9680	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9685	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III
MSG9690	Sensicare PI micro, Surgical glove, Sterile, Synthetic, Polyisoprene, Powder-Free	56293	IIa	III

### \* GMDN code with description

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56293 Polyisoprene surgical glove, non-powdered