

Notification of a Body in the framework of a technical harmonization directive

From : Ministero dello Sviluppo
Economico - Direzione Generale
per il Mercato, la Concorrenza, il
Consumatore, la Vigilanza e la
Normativa Tecnica
Via Sallustiana, 53
00187 ROMA
Italy

To : **European Commission**
GROWTH Directorate-General
200 Rue de la Loi,
B-1049 Brussels.
Other Member States

Reference :

Legislation : 93/42/EEC Medical devices

Body name, address, telephone, fax, email, website :

TUV Rheinland Italia SRL
Via Mattei, 3
20010 - Pogliano Milanese (MI)
Italy
Phone : +39 02 9396871
Fax : +39 02 93968723
Email : informazioni@it.tuv.com
Website : www.tuvitalia.com

Body :

NB 1936

The body is assessed according to :

EN ISO/IEC 17020 - Inspection
EN ISO/IEC 17021 - Certification of management systems
EN ISO/IEC 17065 - Product certification

The competence of the body was assessed by : MINISTERO DELLA SALUTE - MINISTERO DELLO SVILUPPO
ECONOMICO

**The assessment of the body covers the product categories and conformity assessment procedures concerned
by this notification :** Yes

Tasks performed by the Body :

Last approval date : 11/01/2018

| Product family, product /Intended use/Product range | Procedure/Modules | Annexes or articles of the directives | Limitations |
|---|---|---------------------------------------|-------------------------------------|
| *MD 0100 - General non-active, non-implantable medical devices | | | |
| - *MD 0101 - Non-active devices for anaesthesia, emergency and intensive care | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0102 - Non-active devices for injection, infusion, transfusion and dialysis | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | |
| - *MD 0103 - Non-active orthopaedic and rehabilitation devices | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0104 - Non-active medical devices with measuring function | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0105 - Non-active ophthalmologic devices | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0106 - Non-active instruments | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | |
| - *MD 0108 - Non-active medical devices for disinfecting, cleaning, rinsing | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0110 - Non-active medical devices for ingestion | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 0200 - Non-active implants | | | |
| - *MD 0202 - Non-active orthopaedic implants | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | |
| - *MD 0204 - Non-active soft tissue implants | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 0300 - Devices for wound care | | | |

| Product family, product /Intended use/Product range | Procedure/Modules | Annexes or articles of the directives | Limitations |
|--|---|--|--|
| - *MD 0301 - Bandages and wound dressings | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0302 - Suture material and clamps | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | |
| - *MD 0303 - Other medical devices for wound care | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 0400 - Non-active dental devices and accessories | | | |
| - *MD 0401 - Non-active dental equipment and instruments | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0402 - Dental materials | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 0403 - Dental implants | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 1100 - General active medical devices | | | |
| - *MD 1101 - Devices for extra-corporal circulation, infusion and haemopheresis | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1102 - Respiratory devices, devices including hyperbaric chambers for oxygen therapy, inhalation anaesthesia | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices and hyperbaric chambers for oxygen therapy |
| - *MD 1103 - Devices for stimulation or inhibition | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | |
| - *MD 1104 - Active surgical devices | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity | Annex III Annex IV Annex II Annex V | |

| Product family, product /Intended use/Product range | Procedure/Modules | Annexes or articles of the directives | Limitations |
|---|---|--|-------------------------------------|
| | (production quality assurance) EC declaration of conformity (product quality assurance) | Annex VI | |
| - *MD 1105 - Active ophthalmologic devices | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1106 - Active dental devices | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1107 - Active devices for disinfection and sterilisation | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1108 - Active rehabilitation devices and active prostheses | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1109 - Active devices for patient positioning and transport | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1111 - Software | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1112 - Medical gas supply systems and parts thereof | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 1200 - Devices for imaging | | | |
| - *MD 1201 - Imaging devices utilising ionizing radiation | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| *MD 1300 - Monitoring devices | | | |
| - *MD 1301 - Monitoring devices of non-vital physiological parameters | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |

| Product family, product /Intended use/Product range | Procedure/Modules | Annexes or articles of the directives | Limitations |
|---|---|--|-------------------------------------|
| - *MD 1302 - Monitoring devices of vital physiological parameters | EC type-examination EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex III Annex IV Annex II Annex V Annex VI | |
| *MD 1400 - Devices for radiation therapy and thermo therapy | | | |
| - *MD 1401 - Devices utilising ionizing radiation | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1402 - Devices utilising non-ionizing radiation | EC verification EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex IV Annex II Annex V Annex VI | Excluding class III medical devices |
| - *MD 1403 - Devices for hyperthermia / hypothermia | EC declaration of conformity (full quality assurance system) EC declaration of conformity (production quality assurance) EC declaration of conformity (product quality assurance) | Annex II Annex V Annex VI | Excluding class III medical devices |

| Horizontal technical competence | Limitations |
|--|--|
| *MDS 7004 - Medical devices referencing the Directive 2006/42/EC on machinery | |
| *MDS 7006 - Medical devices in sterile condition | Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam) |
| *MDS 7009 - Medical devices utilising biological active coatings and/or materials or being wholly or mainly absorbed | |
| *MDS 7010 - Medical devices incorporating software /utilising software /controlled by software | |