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 (production quality assurance) EC declaration of conformity	Annex II Annex V Annex VI	Excluding class III medical devices
- *MD 0102 - Non-active devices for injection, infusion, transfusion and dialysis	/a	Annex II Annex V Annex VI	
- *MD 0103 - Non-active orthopaedic and rehabilitation devices	EC declaration of conformity	Annex II Annex V Annex VI	Excluding class III medical devices
- *MD 0104 - Non-active medical devices with measuring function	EC declaration of conformity	Annex II Annex V Annex VI	Excluding class III medical devices
- *MD 0105 - Non-active ophthalmologic devices	EC declaration of conformity	Annex II Annex V Annex VI	Excluding class III medical devices
- *MD 0106 - Non-active instruments	EC declaration of conformity	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	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	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	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	Annex III Annex IV Annex II Annex V Annex VI	
- *MD 1104 - Active surgical devices	EC type-examination EC verification 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	Annex IV Annex II Annex V Annex VI	Excluding class III medical devices
- *MD 1403 - Devices for hyperthermia / hypothermia	EC declaration of conformity	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		
	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		