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

From: Office for Registration of Medicinal Products, Medical Devices and Biocidal Products

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To: European Commission

GROWTH Directorate-General

200 Rue de la Loi, B-1049 Brussels.

Other Member States

Reference: Legislation: Regulation (EU) 2017/745 on medical devices

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Body: NB 2274

Tasks performed by the Body :

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Product family, product /Intended use/Product range	Procedure/Modules	Annexes or articles of the directives	Conditions
I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
- A. Active devices - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding MRI
- 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters	Conformity assessment based on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
- 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices for monitoring and/or diagnosis	Conformity assessment based on a quality management		Excluding products used in ophthalmology
3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0302 Active non-implantable devices utilising non-ionizing radiation	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0305 Active non-implantable devices for stimulation or inhibition	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based	Annex IX(I) Annex IX(II) Annex XI(A)	
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0306 Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemapheresis	on a quality management	Annex IX(II)	Including only infusion pumps, devices for dialysis, anaesthesia machines and devices for administration or removal of substances
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	Conformity assessment based on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding hyperbaric chambers
3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0311 Active non-implantable dental devices	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	

Product family, product /Intended use/Product range	Procedure/Modules	Annexes or articles of the directives	Conditions
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport		Annex IX(I) Annex IX(II) Annex XI(A)	Excluding active prostheses
 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	1 112	Annex IX(I) Annex IX(II) Annex XI(A)	
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0316 Medical gas supply systems and parts thereof	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)	
- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation	Conformity assessment based on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
- B. Non-active devices			
- 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	1 105	Annex IX(II) Annex XI(A)	Excluding bone graft substitute for orthopaedic indications, knee, shoulder and hip joint replacement, hyaluronic acid implant for intra-articular use, bone cement
- 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials	Conformity assessment based	Annex IX(I) Annex IX(II) Annex XI(A)	
- 1. Non-active implants and long term surgically invasive devices - MDN 1104 Non-active soft tissue and other implants	Conformity assessment based on a quality management	Annex IX(II) Annex XI(A)	Including only urological tapes, surgical meshes ligament and tendon prostheses made of multifilament polyester fibers
- 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for administration, channelling and removal of substances, including devices for dialysis	Conformity assessment based on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	

Product family, product /Intended use/Product range	Procedure/Modules	Annexes or articles of the directives	Conditions
	Conformity assessment based on product quality assurance		
	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding sutures
	Conformity assessment based on product quality assurance		
- 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
	Conformity assessment based on product quality assurance		
 2. Non-active non-implantable devices MDN 1209 Non-active non-implantable dental materials 	Conformity assessment based	Annex IX(I) Annex IX(II) Annex XI(A)	
	documentation Conformity assessment based on product quality assurance		
2. Non-active non-implantable devices MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based	Annex IX(I) Annex IX(II) Annex XI(A)	
Non-active non-implantable devices MDN 1212 Non-active non-implantable devices for processing and preservation of human cells, tissue or organs including in vitro fertilisation (IVF) and assisted reproductive technologies (ART)	on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based	Annex IX(II)	Excluding devices for in vitro fertilisation (IVF) and assisted reproductive technologies (ART)
- 2. Non-active non-implantable devices - MDN 1213 Non-active non-implantable devices composed of substances to be introduced into the human body via a body orifice or the dermal route	on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	
2. Non-active non-implantable devices MDN 1214 General non-active non-implantable devices used in health care and other non-active non-implantable devices	Conformity assessment based on a quality management	Annex IX(I) Annex IX(II) Annex XI(A)	Including only ultrasound gels, medication cups

Horizontal technical competence	Conditions	
MDS 1001 Devices incorporating medicinal substances		
MDS 1005 Devices in sterile condition	Including: aseptic processing, ethylene oxide gas sterilisation (EOG), moist heat sterilisation, radiation sterilisation, filtration	
MDS 1006 Reusable surgical instruments		
MDS 1007 Devices incorporating or consisting of nanomaterial		
MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly		

Horizontal technical competence	Conditions	
or mainly absorbed or locally dispersed in the human body or are intended to undergo a chemical change in the body		
MDS 1009 Devices incorporating software/utilising software/controlled by software, including devices intended for controlling, monitoring or directly influencing the performance of active or active implantable devices		
MDS 1010 Devices with a measuring function		
MDS 1011 Devices in systems or procedure packs		
MDS 1012 Products without an intended medical purpose listed in Annex XVI to Regulation (EU) 2017/745	Limited to devices emitting electromagnetic radiation for use on the human body according to Annex XVI p. 5	
MDS 1013 Class III custom-made implantable devices		
MDT 2001 Devices manufactured using metal processing		
MDT 2002 Devices manufactured using plastic processing		
MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)	Excluding processing of glass	
MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)	Excluding processing of leather	
MDT 2006 Devices manufactured using chemical processing		
MDT 2007 Devices which require knowledge regarding the production of pharmaceuticals		
MDT 2008 Devices manufactured in clean rooms and associated controlled environments		
MDT 2009 Devices manufactured using processing of materials of human, animal, or microbial origin	Excluding processing of animal materials	
MDT 2010 Devices manufactured using electronic components including communication devices		
MDT 2011 Devices which require packaging, including labelling		
MDT 2012 Devices which require installation, refurbishment	Limited to electronic devices and medical gas installations	