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

From:

Swedish Medical Products Agency **To:** Box 26, SE-751 03 Dag Hammarskjölds väg 42, SE-752 37 Uppsala Sweden 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

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

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# **Body info:**

NB 3033

#### Body:

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## Tasks performed by the Body:

Last approval date: 2024-03-07

Products	Procedures	Articles /Annexes	Conditions
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -2. Active non-implantable devices		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded, EEG excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -2. Active non-implantable devices	<b>o</b> ,	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded, Short wave diathermy excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(II) Annex XI(A)	Class III Excluded, Heart-lung-bypass pump/Extracorpore al Membrane Oxygenation (ECMO) excluded, Intra-aorthic Balloon Pump excluded

Products	Procedures	Articles /Annexes	Conditions
-I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable therapeutic devices and general active non-implantable devices -MDA 0307 Active non- implantable respiratory devices	<b>3</b>	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
-I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable therapeutic devices and general active non-implantable devices -MDA 0308 Active non- implantable devices for wound and skin care		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
-I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable therapeutic devices and general active non-implantable devices -MDA 0310 Active non- implantable devices for ear, nose and throat	<b>3</b>	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
-I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable therapeutic devices and general active non-implantable devices -MDA 0311 Active non- implantable dental devices		Annex IX(II) Annex XI(A)	Class III Excluded, Restricted to: Powered dental surgical unit and hand pieces, Surgical suction device for dental use
<ul> <li>-I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE</li> <li>-A. Active devices</li> <li>-3. Active non-implantable therapeutic devices and general active non-implantable devices</li> <li>-MDA 0312 Other active non- implantable surgical devices</li> </ul>		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded

Products	Procedures	Articles /Annexes	Conditions
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded, Medical gas pipeline system excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded, Sterilization devices excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -A. Active devices -3. Active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded

Products	Procedures	Articles /Annexes	Conditions
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -1. Non-active implants and long	<b>o</b> ,	Annex IX(II) Annex XI(A)	Class III Excluded, Restricted to: Orthopaedic nails, screws, plates, and Sutures, suture anchors, staples for orthopedic surgery
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -1. Non-active implants and long		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable	<b>o</b> ,	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable	<b>č</b>	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded, Restricted to: Periferal vascular catheters and related tools

Products	Procedures	Articles /Annexes	Conditions
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable	<b>o</b> ,	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE		Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable	<b>S 1</b>	Annex IX(II) Annex XI(A)	Class III Excluded, Catheter lock solutions excluded, and Solutions for disinfecting contact lenses excluded
DESIGN AND INTENDED PURPOSE OF THE DEVICE -B. Non-active devices -2. Non-active non-implantable	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)	Class III Excluded

Но	prizontal technical competences	Limitations
	Devices which are also machinery as (a) of the second paragraph of Article 2 of 2/EC of the European Parliament and of	
MDS 1005	Devices in sterile condition:	Restricted to: ethylene oxide gas sterilisation (EOG) and radiation sterilisation (gamma, x-ray, electron beam)
MDS 1006	Reusable surgical instruments:	
intended for cor	Devices incorporating software/utilising lled by software, including devices ntrolling, monitoring or directly influencing e of active or active implantable devices: Devices with a measuring function:	
MDS 1011	Devices in systems or procedure packs:	
MDT 2001 processing:	Devices manufactured using metal	
MDT 2002 processing:	Devices manufactured using plastic	
MDT 2003 mineral process	Devices manufactured using non-metal sing (e.g. glass, ceramics):	
MDT 2004 non-mineral pro paper):	Devices manufactured using non-metal ocessing (e.g. textiles, rubber, leather,	
MDT 2006 processing:	Devices manufactured using chemical	
MDT 2008 and associated	Devices manufactured in clean rooms controlled environments:	
MDT 2010 components inc	Devices manufactured using electronic cluding communication devices:	
MDT 2011 including labelli	Devices which require packaging, ing:	
MDT 2012 refurbishment:	Devices which require installation,	