Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
ÜV NORD CERT GmbH ım TÜV 1 5307 Essen Germany	0044	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
Germany		 - A. Active devices - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)	
		- 2. Active non-implantable devices for imaging, monitoring and/or diagnosis	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)	
		- Z. Active non-implantable devices for imaging.	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)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			Conformity assessment based on product quality assurance						
		 - 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 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 0301 Active non-implantable devices utilising ionizing radiation 	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 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF B Name and address of the notified	NOTIFIED UNDER DIRECTIVE : Regulat Responsible for the following products	Responsible for the	Annexes or	es Conditions
bodies	/Horizontal technical competence	following procedures or modules	articles of the directives	
		based on product quality assurance		
	 - 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	Annex IX(I) Annex IX(II) Annex XI(A)	
		based on product quality assurance		
	 - 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 	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 0307 Active non-implantable respiratory devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	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 0309 Active non-implantable ophthalmologic devices 	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 0310 Active non-implantable devices for ear, nose and throat 		Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices	Conformity assessment based on a quality	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 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		-		Annex IX(I)				

LIST OF E	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical devi	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		general active non-implantable devices - MDA 0315 Software	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)	
		- 3. Active non-implantable therapeutic devices and	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 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)	
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0318 Other active non-implantable devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)			
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)			
		-	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	7		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)				
		- - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- MDN 1203 Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
		- 2. Non-active non-implantable devices	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)				
		- 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex IX(I)				
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			technical documentation Conformity assessment based on product quality assurance				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)			
		- 2. Non-active non-implantable devices	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)			
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		 - 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 	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)	
		- - 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 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)	
		MDS 1001 Devices incorporating medicinal substances			without medicinal products derived from human blood or human plasma
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		MDS 1005 Devices in sterile condition			including: aseptic processing; ethylene oxide gas sterilisation (EOG); low temperature steam and formaldehyde sterilisation; moist heat sterilisation; radiation sterilisation (gamma, x-ray, electron beam); sterilisation with hydrogen peroxide; additional: Sterilisationsverfahren mit Plasma			
		MDS 1006 Reusable surgical instruments						
		MDS 1007 Devices incorporating or consisting of nanomaterial						
		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 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather,						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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 2010 Devices manufactured using electronic components including communication devices					
		MDT 2011 Devices which require packaging, including labelling					
		MDT 2012 Devices which require installation, refurbishment					
		MDT 2013 Devices which have undergone reprocessing					
ational Standards Authority of Ireland (NSAI) Swift Square, Northwood, Santry ublin 9 eland	0050	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- B. Non-active devices					
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			
			based on product quality				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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)				
			Conformity assessment based on product quality assurance					
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	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	assurance 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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)				
		 - - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDN 1204 Non-active non-implantable devices for wound and skin care	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)			
		- 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)			
		- 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)			
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		- 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)			
		- 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental	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)			
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		disinfecting, cleaning and rinsing	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance						
		- - 2. 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)	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)					
		 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 	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)					
		 - 2. Non-active non-implantable devices - MDN 1214 General non-active non-implantable devices used in health care and other non-active 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		non-implantable devices	based on assessment of technical documentation Conformity assessment based on product quality assurance				
		- A. Active devices					
		- - 1. Active implantable devices - MDA 0101 Active implantable devices for stimulation/inhibition/monitoring	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)			
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)			
		- - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters	based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 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 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 0309 Active non-implantable ophthalmologic devices 	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 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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 0315 Software 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 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 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 0318 Other active non-implantable devices 	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 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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 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 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 0306 Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemapheresis 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			assurance						
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	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)					
		MDS 1001 Devices incorporating medicinal substances							
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives							
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)							
		MDS 1005 Devices in sterile condition			Including EtO, Moist Heat, Asepti Chemical, Irradiation				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		MDS 1006 Reusable surgical instruments						
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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						
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		MDT 2006 Devices manufactured using chemical						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
IMQ ISTITUTO ITALIANO DEL MARCHIO DI QUALITÀ S.P.A. Via Quintiliano, 43 20138 - MILANO Italy	0051	 1. Active implantable devices MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	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)				
			Conformity assessment based on product quality assurance					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical device	S
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	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 0316 Medical gas supply systems and parts thereof	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		 - - 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 type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on assessment of technical documentation		
			Conformity assessment based on product quality assurance		
			Conformity assessment based on product conformity verification		
		3. Active non-implantable therapeutic devices and	Conformity assessment based on type-examination	Annex X Annex IX(I)	
		general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(II) Annex XI(A)	
			Conformity assessment based on assessment of technical documentation	Annex XI(B)	
			Conformity assessment based on product quality assurance		
			Conformity assessment based on product conformity verification		
		- B. Non-active devices			
		1. Non-active implants and long term surgically invasive devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III devices
			Conformity assessment		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1102 Non-active osteo- and orthopaedic implants	based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials	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)	Excluding class III devices			
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)	Excluding class III devices			
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)	Excluding class III devices			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)	Excluding class III devices			
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	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 class III devices			

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			Conformity assessment based on product quality assurance						
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)	Excluding class III devices				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	Excluding class III devices				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices fo disinfecting, cleaning and rinsing	Conformity assessment based on a quality management system r Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III devices				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		assisted reproductive technologies (ART)	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(II) Annex XI(A)	Excluding class III devices			
		 - 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 	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)	Excluding class III devices			
		- 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III devices			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	assurance 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)	
		 - 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 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)	Excluding class III devices
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Excluding formaldehyde sterilization.
		MDS 1006 Reusable surgical instruments			
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly or mainly absorbed or			

ame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2006 Devices manufactured using chemical processing			
		MDT 2008 Devices manufactured in clean rooms and associated controlled environments			
		MDT 2010 Devices manufactured using electronic components including communication devices			
		MDT 2011 Devices which require packaging, including			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
ÜV SÜD Product Service GmbH Lidlerstraße 65 0339 MÜNCHEN Germany	0123	labelling MDT 2012 Devices which require installation, refurbishment MDT 2013 Devices which have undergone reprocessing I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE						
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product conformity verification					
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials 	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)				
			Conformity assessment based on product quality assurance					
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1104 Non-active soft tissue and other implants 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation		For breast implants only Annex IX applicable			
			Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification					
		- - 2. Non-active non-implantable devices	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
			Conformity assessment					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product conformity verification					
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)				
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	assurance Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on product quality assurance				
			Conformity assessment based on product conformity verification				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			
			based on product quality assurance				
		Instruments	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)			
		 - - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on assessment of technical documentation Conformity assessment based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)	
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)	
		 - 2. 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 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		assisted reproductive technologies (ART)	technical documentation Conformity assessment based on product quality assurance			
		 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 	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)		
		- - 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 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)		
		- A. Active devices	Conformity assessment	Annex X		
		 1. Active implantable devices MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	based on type-examination Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)			
		- - 1. Active implantable devices - MDA 0102 Active implantable devices delivering drugs or other substances	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 1. Active implantable devices - MDA 0103 Active implantable devices supporting or replacing organ functions 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification				
		- 1. Active implantable devices - MDA 0104 Active implantable devices utilising radiation and other active implantable devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	based on product conformity verification Conformity assessment	Annex X Annex IX(I)			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices 	Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 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 type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives			
		- - 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 assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination	Annex X Annex IX(I)			
		- MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	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(II) Annex XI(A) Annex XI(B)			
			Conformity assessment based on product conformity verification				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for 	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0305 Active non-implantable devices for	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		of substances and haemapheresis	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
			based on product conformity verification Conformity assessment	Annex X			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear. 		Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(I) Annex XI(A) Annex XI(B)			
			Conformity assessment based on product conformity verification				
		general delive nen implantable deviced	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- MDA 0313 Active non-implantable prostneses.	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues 	Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		assisted reproductive technologies (ART)	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	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 0316 Medical gas supply systems and parts thereof	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance Conformity assessment based on product conformity verification					
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			assurance Conformity assessment based on product conformity verification		
		 - 3. Active non-implantable inerapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		MDS 1001 Devices incorporating medicinal substances	-		
		MDS 1002 Devices manufactured utilising tissues or cells of human origin, or their derivatives			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Including aseptic processing,

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
					ethylene oxide gas sterilisation (EOG), low temperature steam and formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), sterilisation with hydrogen peroxide, sterilisation with liquid chemical sterilising agents, thermic sterilisation with dry heat				
		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 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							
		MDS 1013 Class III custom-made implantable devices							
		MDS 1014 Devices incorporating as an integral part an							

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		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						
		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						
		MDT 2013 Devices which have undergone reprocessing			Only for medical devices that ar forseen by the manufacturer to			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
					undergo reprcessing			
DEKRA Certification GmbH landwerkstraße 15 0565 STUTTGART Germany	0124	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE						
		- A. Active devices						
		- 2. Active non-implantable devices for imaging,	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)				
	- MDA 0	- MDA 0201 Active non-implantable imaging devices	Conformity assessment	Annex XI(A)				
			Conformity assessment based on product quality assurance					
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		- MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation	based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		- 2. Active non-implantable devices for imaging.	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
			Conformity assessment based on assessment of					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		monitoring of vital physiological parameters	technical documentation Conformity assessment based on product quality assurance					
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices 	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 0301 Active non-implantable devices utilising ionizing radiation 	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 0302 Active non-implantable devices utilising non-ionizing radiation 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 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 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	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	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 	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 0307 Active non-implantable respiratory devices 	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 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- 5. Active non-implantable therapeutic devices and	assurance 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 0311 Active non-implantable dental devices	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)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	.,	excluding IVF and ART; limited to devices for cryopreservation
		-	Conformity assessment	Annex IX(I)	

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Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0318 Other active non-implantable devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)				
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)				
		-	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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)	excluding breast implants whose purpose is the enlargement or replacement of the volume of the breast			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)				
		- - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

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Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		administration, channelling and removal of substances, including devices for dialysis	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on assessment of technical documentation Conformity assessment					
			based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				
			based on product quality assurance					
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	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)				
			Conformity assessment based on product quality assurance					
		 - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		 - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 2. 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) 	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)				
		- 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	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)	excluding devices for ingestion			
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1002 Devices manufactured utilising tissues or cells of human origin, or their derivatives			restricted to devices manufactured utilising human serum albumin (HSA)			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			including aseptic processing; ethylene oxide gas sterilisation (EOG); low temperature steam and formaldehyde sterilisation; moist heat sterilisation; radiation sterilisation (gamma, x-ray, electron beam); sterilisation with hydrogen peroxide; sterilisation with liquid chemical sterilising agens;			
		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 or mainly absorbed or locally dispersed in the human body or are intended to						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		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			only products within the designation according to MDA/MDN-codes excluding products according to section 3 of annex XVI		
		MDS 1013 Class III custom-made implantable devices			only products within the designation according to MDA/MDN-codes		
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device					
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		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			only products within the designation according to MDA/MDN-codes		
		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					
		MDT 2013 Devices which have undergone reprocessing			excluding reprocessing of single-use devices		
ΓÜV Rheinland LGA Products GmbH Fillystraße 2 90431 Nürnberg Germany	0197	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- A. Active devices					
		 - 1. Active implantable devices - MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	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)	limited to stimulation devices excluding brain stimulators and pacemakers		

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		- 2. Active non-implantable devices for imaging.	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)	
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
			based on product quality assurance Conformity assessment based on product conformity verification		
			Conformity assessment based on type-examination	Annex X Annex IX(I)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		monitoring and/or diagnosis - MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(II) Annex XI(A) Annex XI(B)				
			based on product conformity verification					
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy)	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	excluding brain stimulation devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	excluding hyperbaric chamber			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART)	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation 	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product conformity verification		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(I) Annex XI(A) Annex XI(B)	
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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(II)	excluding heart valves introduce into the body by open heart surgeries
		-	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	excluding joint implants

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Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)				
		- - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

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Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 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 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)	
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)	
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on assessment of technical documentation Conformity assessment based on product quality						
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	assurance 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)					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)					
		 - - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		 - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases 	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)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical devic	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			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 	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)	
		 - 2. Non-active non-implantable devices - MDN 1212 Non-active non-implantable devices for processing and preservation of human cells, tissue 	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)	
		 - 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 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- 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 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)				
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			including aseptic processing ethylene oxide gas sterilisation (EOG) low temperature steam and formaldehyde sterilisation moist heat sterilisation radiation sterilisation (gamma, x-ray, electron beam) sterilisation with hydrogen peroxide sterilisation with liquid chemical sterilising agens thermic sterilisation with dry heat			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		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 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						
		MDS 1013 Class III custom-made implantable devices						
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather,						

LIST OF E	ODIES	NOTIFIED UNDER DIRECTIVE : Regulati	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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 "under processing of materials of human origin"
		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			
		MDT 2013 Devices which have undergone reprocessing			
DQS Medizinprodukte GmbH August-Schanz-Straße 21 60433 FRANKFURT AM MAIN Germany	0297	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
<i>.</i>		- A. Active devices			
		 - 1. Active implantable devices - MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(II)	restricted to active implantable devices for cardiovascular/vascula stimulation / inhibition / monitoring

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 2. Active non-implantable devices for imaging, monitoring and/or diagnosis MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)				
		 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)				
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 o	n medical device	25
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)	
		- MDA 0301 Active non-implantable devices utilising ionizing radiation	Conformity assessment based on assessment of technical documentation		
		- 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	
		non-ionizing radiation	Conformity assessment based on assessment of technical documentation Conformity assessment		
			based on product quality assurance		
		- 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II)	
		shock-wave therapy (lithotripsy)	technical documentation Conformity assessment	Annex IX(I)	
		I - 3. Active non-implantable therapeutic devices and	based on a quality management system	Annex IX(II) Annex XI(A)	
		- MDA 0305 Active non-implantable devices for stimulation or inhibition	Conformity assessment based on assessment of		

ame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 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 		Annex IX(I) Annex IX(II) Annex XI(A)	
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices 	Conformity assessment	Annex IX(I) Annex IX(II)	
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	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 0310 Active non-implantable devices for ear, nose and throat 	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)	restricted to external hearing aids			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 	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(II)	restricted to active non-implantable devices for patient positioning and transport		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)			

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		general active non-implantable devices - MDA 0316 Medical gas supply systems and parts thereof	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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)					
		 B. Non-active devices - 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)					
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		implants	technical documentation				
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)			
		 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)			
		- - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for	assurance Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		administration, channelling and removal of substances, including devices for dialysis	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on assessment of technical documentation						
			Conformity assessment based on product quality assurance						
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex IX(I)					
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			technical documentation Conformity assessment based on product quality assurance				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)			
		2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)			
		 - 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 	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)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 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 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)				
		MDS 1001 Devices incorporating medicinal substances MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			including aseptic processing; ethylene oxide gas sterilisation (EOG); low temperature steam and formaldehyde sterilisation; moist heat sterilisation; radiation sterilisation (gamma, x-ray, electron beam); sterilisation with hydrogen peroxide; thermic sterilisation with dry heat			

LIST OF B	ODIE	S NOTIFIED UNDER DIRECTIVE : Regulation	on (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		MDS 1006 Reusable surgical instruments			
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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			restricted to products corresponding Regulation (EU) 2017/745 in Annex XVI section 1. and section 2.
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		MDT 2006 Devices manufactured using chemical processing			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		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			restricted to devices manufactured using processing of materials of animal or microbial origin			
		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						
CENTRO NACIONAL DE CERTIFICACION DE PRODUCTOS SANITARIOS Campezo 1. Edificio 7. 28022 MADRID Spain	0318	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 0201 Active non-implantable imaging devices utilising ionizing radiation 	based on assessment of technical documentation Conformity assessment based on product quality		Limited to x-ray medical devices, gamma cameras and positron emission tomography			
			assurance Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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 0302 Active non-implantable devices utilising non-ionizing radiation 	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)	Limited to medical devices for magnetotherapy and microwaves			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	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)	Limited to diagnostic medical devices		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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 0315 Software 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0316 Medical gas supply systems and parts 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices				
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on product quality assurance						
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)					
		- B. Non-active devices	assurance						
						- - 1. Non-active implants and long term surgically invasive devices - MDN 1101 Non-active cardiovascular, vascular	Conformity assessment based on a quality management system Conformity assessment	Annex IX(II) cardiovascular surgery, Annex XI(A) implantable drug deliver	Limited to stents, sutures for cardiovascular surgery, and implantable drug delivery systems
		and neurovascular implants	based on assessment of technical documentation Conformity assessment based on product quality assurance						
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)				
		 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		Neurological, neurosurgical and breast implants are excluded			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			assurance		
		- 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 type-examination 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 X limited to medical devices for puncture, injection and/or extraction of fluids
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)	
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
			Conformity assessment based on product quality assurance					
		 - 2. Non-active non-implantable devices MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases 	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)				
			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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system		Annex X limited to contact lens care products			
			Conformity assessment based on assessment of technical documentation Conformity assessment					

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on product quality assurance						
		 - 2. 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) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex IX(I)					
		- 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	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance				
		MDS 1001 Devices incorporating medicinal substances					
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			Excluding medical devices utilising tissues or cells of animal origin under Regulation (UE) No. 722/2012		
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)					
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam and formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam)		
		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 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					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		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				
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device				
		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)				
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)				
		MDT 2005 Devices manufactured using biotechnology				
		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 human and animal material of Regulation (UE) No. 722/2012	
		MDT 2010 Devices manufactured using electronic				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
DEKRA Certification B.V. Meander 1051 / P.O. Box 5185 5825 MJ ARNHEM / 6802 ED ARNHEM Netherlands	0344	 1. Active implantable devices MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(I) Annex XI(A) Annex XI(B)			
		- - 1. Active implantable devices	Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I)			
		- MDA 0102 Active implantable devices delivering	Comonnity assessment	Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		drugs or other substances	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)			
		- - 1. Active implantable devices - MDA 0103 Active implantable devices supporting or replacing organ functions	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 1. Active implantable devices - MDA 0104 Active implantable devices utilising	Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 2. Active non-implantable devices for imaging,	Conformity assessment	Annex X Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
				Annex XI(A) Annex XI(B)			
		 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 system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 2. Active non-implantable devices for imaging,	Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		for monitoring and/or diagnosis	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
				Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy)	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)			
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		extra-corporal circulation, administration or removal of substances and haemapheresis	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)				
		 - 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 system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- 3. Active non-implantable therapeutic devices and		Annex X Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0308 Active non-implantable devices for wound and skin care	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)			
		devices	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0310 Active non-implantable devices for ear, nose and throat	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)				
		 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0311 Active non-implantable dental devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0312 Other active non-implantable surgical devices	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)				
		 - 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 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II)				

			1		
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0315 Software	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment	Annex XI(A) Annex XI(B) Annex X	
		 - 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	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I) Annex IX(II)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- B. Non-active devices	Conformity assessment	Annex X			
		-	based on type-examination	Annex IX(I)			

LIST OF E	ODIES	NOTIFIED UNDER DIRECTIVE : Regula	ation (EU) 2017/745 or	n medical devi	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(II) Annex XI(A) Annex XI(B)	
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		1. Non-active implants and long term surgically	Conformity assessment based on type-examination	Annex X Annex IX(I)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		invasive devices - MDN 1103 Non-active dental implants and dental materials	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 1. Non-active implants and long term surgically	based on product conformity verification Conformity assessment based on type-examination	Annex X Annex IX(I)				
		invasive devices - MDN 1104 Non-active soft tissue and other implants	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(II) Annex XI(A) Annex XI(B)				
			Conformity assessment based on product quality assurance Conformity assessment					
			based on product conformity verification Conformity assessment	Annex X				
		- - 2. Non-active non-implantable devices	based on type-examination	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(II) Annex XI(A) Annex XI(B)			
			based on product conformity verification Conformity assessment				
		 - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for administration, channelling and removal of substances, including devices for dialysis 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
			Conformity assessment based on product quality assurance Conformity assessment based on product				
		-	conformity verification Conformity assessment based on type-examination	Annex X Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		introducers, filters, and related tools	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(II) Annex XI(A) Annex XI(B)				
			based on product quality assurance Conformity assessment based on product conformity verification	Annex X ⁿ Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of					
			technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product					
		-	conformity verification Conformity assessment based on type-examination	Annex X Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDN 1206 Non-active non-implantable ophthalmologic devices	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(II) Annex XI(A) Annex XI(B)			
			assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification				
		- - 2. Non-active non-implantable devices	Conformity assessment based on type-examination	Annex X Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDN 1208 Non-active non-implantable instruments	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(II) Annex XI(A) Annex XI(B)			
			Conformity assessment based on product conformity verification	Annex X ⁿ Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification				
		- - 2. Non-active non-implantable devices	Conformity assessment based on type-examination	Annex X Annex IX(I)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A) Annex XI(B)	
	Conformity assessment based on product conformity verification				
		 - 2. 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
	Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification				
			Conformity assessment based on type-examination	Annex X Annex IX(I)	

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 oi	n medical dev	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- 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	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(II) Annex XI(A) Annex XI(B)	
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		MDS 1001 Devices incorporating medicinal substances			
		MDS 1002 Devices manufactured utilising tissues or			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulation	on (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		cells of human origin, or their derivatives			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			including aseptic processing, ethylene oxide gas sterilization (EOG), low temperature steam and formaldehyde sterilization, moist heat sterilization, radiation sterilization (gamma, x-ray, electron beam), sterilisation with hydrogen peroxide, sterilization with liquid chemical sterilizing agents, thermic sterilization with dry heat
		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 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			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical device	25
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			
		MDS 1013 Class III custom-made implantable devices			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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			
		MDT 2010 Devices manufactured using electronic			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		components including communication devices				
		MDT 2011 Devices which require packaging, including labelling				
		MDT 2012 Devices which require installation, refurbishment				
		MDT 2013 Devices which have undergone reprocessing				
STITUTO SUPERIORE DI SANITA' 'iale Regina Elena, 299 0161 - ROMA aly	0373	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 0301 Active non-implantable devices utilising 	Conformity assessment based on type-examination Conformity assessment based on product conformity verification	Annex X Annex XI(B)	Limited to therapeutic cyclotrons and linear accelerators	
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)		

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical devic	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0305 Active non-implantable devices for stimulation or inhibition	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	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 0310 Active non-implantable devices for ear, 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		nose and throat	based on assessment of technical documentation Conformity assessment based on product quality assurance			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)		

- - Conformity assessment based on product quality assessment based on a quality management system Annex IX(I) - - Conformity assessment based on a quality management system Annex IX(I) - - Conformity assessment based on a quality management system Annex IX(I) - - Conformity assessment based on a sessessment based on a sessessment based on a sessessment based on product quality assessment based on product quality assessment based on a quality management system Annex IX(I) - - - Conformity assessment based on a sessessment based on a sessessment based on a quality assessment based on a sessessment of technical documentation Conformity assessment based on assessment to assessment to assed on a product quality assessment based on assessment to active non-implantable devices Annex IX(I) - - - - - - - - - - - - - - - - - - -	Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
- 3. Active non-implantable therapeutic devices - MDA 0316 Medical gas supply systems and parts thereof based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Annex IX(II) Annex XI(A) - 3. Active non-implantable therapeutic devices an general active non-implantable therapeutic devices - MDA 0318 Other active non-implantable devices Conformity assessment based on a quality management system Annex IX(II) Annex XI(A) - MDA 0318 Other active non-implantable devices - MDA 0318 Other active non-implantable devices Conformity assessment based on a quality management system Annex IX(I) Annex XI(A) - B. Non-active devices - Conformity assessment based on a quality Annex IX(I) - B. Non-active devices - Conformity assessment based on a quality Annex IX(I) - B. Non-active devices - Conformity assessment based on a quality Annex IX(I)				Conformity assessment based on product quality		
 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices - Conformity assessment based on product quality assurance - B. Non-active devices - Conformity assessment - B. Non-active devices - Conformity assessment - Non-active devices -			 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0316 Medical gas supply systems and parts thereof 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	
- Conformity assessment Annex IX(I)			 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	
AIMEX IX.			-	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 or	n medical device	S
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		and neurovascular implants	technical documentation Conformity assessment based on product quality assurance		
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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)	
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)	
		 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical devi	ces
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		 - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care 	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)	
		 - 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 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)	
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical devic	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)	
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical devi	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			assurance		
		- 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)	
		- 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)	
		- 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
		- - 2. 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)	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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)				
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam, moist heat sterilisation, radiation			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
					sterilisation (gamma-ray, electron beam), dry heat.		
		MDS 1006 Reusable surgical instruments					
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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					
		MDS 1013 Class III custom-made implantable devices					
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device					
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather,					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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					
		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					
CIM S.P.A. Piazza Don Enrico Mapelli, 75 20099 - Sesto San Giovanni (MI) taly	0425	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 0203 Active non-implantable devices for monitoring of vital physiological parameters 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 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 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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices			
		 - - 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 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex IX(I)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- B. Non-active devices	assurance					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)	Excluded Class III Medical Devices			
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1202 Non-active non-implantable devices for administration, channelling and removal of substances, including devices for dialysis	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)				
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		instruments	Conformity assessment based on assessment of technical documentation		
			Conformity assessment based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)	
		 - 2. Non-active non-implantable devices - MDN 1213 Non-active non-implantable devices composed of substances to be introduced into the 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		human body via a body orifice or the dermal route	based on assessment of technical documentation Conformity assessment based on product quality assurance					
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilisation (EOG) - moist heat sterilisation - radiation sterilisation (gamma-ray x-ray, electron beam)			
		MDS 1006 Reusable surgical instruments						
		MDS 1007 Devices incorporating or consisting of nanomaterial						
		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						
		MDT 2001 Devices manufactured using metal processing						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		MDT 2002 Devices manufactured using plastic processing					
		MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		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 2010 Devices manufactured using electronic components including communication devices					
		MDT 2011 Devices which require packaging, including labelling					
		MDT 2012 Devices which require installation, refurbishment					
ITALCERT SRL Viale Sarca, 336 20126 - MILANO Italy	0426	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- A. Active devices					
		2. Active non-implantable devices for imaging,	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)			
		monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices	Conformity assessment	Annex XI(A)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical devic	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		for monitoring and/or diagnosis	based on assessment of technical documentation Conformity assessment based on product quality assurance		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	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 	assurance 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)	
		 B. Non-active devices - 1. Non-active implants and long term surgically invasive devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1102 Non-active osteo- and orthopaedic implants	based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials	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)				
		- - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical device	S
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
		 - 2. 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) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	Excluded devices for In Vitro Fertilisation (IVF) and Assisted Reproductive Technologies (ART)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 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 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				
			based on product quality assurance					
		MDS 1001 Devices incorporating medicinal substances			Excluded human blood or plasma derivatives			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			Excluded medical devices utilising tissues of animal origin under Commission Reulation (UE) No. 722/2012.			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilisation (EOG) - moist heat sterilisation - radiation sterilisation (gamma-ray, x-ray, electron beam) - hydrogen peroxide sterilisation			
		MDS 1006 Reusable surgical instruments						
		MDS 1007 Devices incorporating or consisting of						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		nanomaterial					
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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					
		MDS 1013 Class III custom-made implantable devices			Limited to non-active osteo- and orthopaedic implants, non-active dental implants and dental materials, non-active soft tissue and other implants		
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		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						
		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						
GMED SAS 1, rue Gaston Boissier 75015 PARIS France	0459	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE						
		- B. Non-active devices						
			Conformity assessment based on type-examination	Annex X Annex IX(I)				
		- MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system	Annex IX(II) Annex XI(A)				
			Conformity assessment based on assessment of technical documentation					
			Conformity assessment					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on product quality assurance				
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable 	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance Conformity assessment	Annex X			
		- 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	based on type-examination 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)			
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		 - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		 - 2. 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) 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 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	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	assurance Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		-	Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	based on type-examination 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)			
		 - A. Active devices - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			
		-	Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation	Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
			conformity verification Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0305 Active non-implantable devices for stimulation or inhibition	Conformity assessment based on a quality	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 S. 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 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	conformity verification Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		-	conformity verification Conformity assessment	Annex X				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0316 Medical gas supply systems and parts thereof 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	conformity verification Conformity assessment	Annex X			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Active implantable devices MDA 0104 Active implantable devices utilising radiation and other active implantable devices 	based on type-examination 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)				
		- - 1. Active implantable devices - MDA 0101 Active implantable devices for stimulation/inhibition/monitoring	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 1. Active implantable devices - MDA 0102 Active implantable devices delivering drugs or other substances	assurance Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		- - 1. Active implantable devices - MDA 0103 Active implantable devices supporting or replacing organ functions	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex X Annex IX(I) Annex IX(II) Annex XI(A)	
		MDS 1001 Devices incorporating medicinal substances			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			The covered sterilization proces are : - aseptic processing - ethylene oxide gas sterilisation (EOG), - low temperature stean formaldehyde sterilization, - mo heat sterilization, - radiation

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
					sterilisation (gamma, x-ray, electron beam), - hydrogen peroxyde, - liquid chemical sterilising agents, - dry heat.
		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 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			
		MDS 1013 Class III custom-made implantable devices			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		MDT 2001 Devices manufactured using metal processing			
		MDT 2002 Devices manufactured using plastic processing			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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			
		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			
		MDT 2013 Devices which have undergone reprocessing			
IWA CERMET ITALIA S.P.A. l'a Cadriano, 23 0057 - Cadriano di Granarolo (BO) aly	0476	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
		- B. Non-active devices			
		-	Conformity assessment	Annex IX(I)	
		- 2. Non-active non-implantable devices	based on a quality	Annex IX(II)	

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Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 - 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 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)			
		- 2. Non-active non-implantable devices	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)			
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		wound and skin care	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality				
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	assurance 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)			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)			
		 - - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on assessment of technical documentation Conformity assessment based on product quality assurance						
		- 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)					
		 - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing 	assurance 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)					
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(II)	Excluded in vitro fertilisation (IVF) and assisted reproductive technologies (ART)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		assisted reproductive technologies (ART)	technical documentation Conformity assessment based on product quality assurance					
		 - 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 	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)				
		- - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)				
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials 	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)				
		 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- A. Active devices						
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				
			Conformity assessment based on product quality assurance					
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded magnetic resonance			
		 MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
	- MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters	Conformity assessment based on assessment of technical documentation						
			Conformity assessment					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 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 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 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 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 0306 Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemapheresis 	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 0307 Active non-implantable respiratory devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	.,	Excluding hyperbaric chamber for oxygen therapy			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	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 0309 Active non-implantable ophthalmologic devices 	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 0310 Active non-implantable devices for ear, nose and throat 	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)	Limited to ear equipment			
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0311 Active non-implantable dental devices	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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)				
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Excluded in vitro fertilisation (IVF) and assisted reproductive			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART)	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)	technologies (ART)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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 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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)					

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	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)	
		MDS 1001 Devices incorporating medicinal substances			Excluding plasma and blood derivatives
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			The following processes are covered: aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam formaldehyde sterilisation, mois

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
					heat sterilisation, dry heat sterilization, radiation sterilisation (gamma-ray, x-ray, electron beam)		
		MDS 1006 Reusable surgical instruments					
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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					
		MDS 1013 Class III custom-made implantable devices			Limited to devices within the scope of designation related to MDN codes		
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device					
		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)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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			Limited to processing of materials of animal and microbial origin		
		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 devices which require installation		
		MDT 2013 Devices which have undergone reprocessing			Limited to reusable devices which have to undergone reprocessing		
Eurofins Product Testing Italy S.r.l. /ia Courgnè, 21 I0156 - TORINO (TO) taly	0477	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- A. Active devices					
		- 2. Active non-implantable devices for imaging.	Conformity assessment based on a quality	Annex IX(I) Annex XI(A)	Excluded Class III Medical Device		
		monitoring and/or diagnosis	management system	Annex XI(B)			

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance Conformity assessment					
			based on product conformity verification					
		 - 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 system Conformity assessment based on product quality assurance Conformity assessment based on product	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	based on a quality	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			
		- 3. Active non-implantable therapeutic devices and general active non-implantable devices		Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		hyperthermia/hypothermia	based on product quality assurance Conformity assessment based on product conformity verification					
		 - 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 product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			
		 - 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 system Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on product quality	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance Conformity assessment based on product conformity verification				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance Conformity assessment based on product	Annex XI(A)	Excluded Class III Medical Devices Excluded active non implantable devices for nose and throat		
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	conformity verification Conformity assessment based on a quality management system Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity verification Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Excluded Class III Medical Devices		

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 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 system Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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)	Excluded Class III Medical Devices excepted those classified in Class III only composed of substances or a combination of substances that are absorbed by or locally dispersed in the human body and/or utilising tissues of animal origin, including Commission Regulation (UE) n. 722/2012.
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on product quality assurance Conformity assessment based on product conformity verification				
		 - 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 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)			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)	Excluded Class III Medical Devices		
		 - 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 type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices		

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification					
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 2. Non-active non-implantable devices MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	Conformity assessment based on a quality management system Conformity assessment based on product quality	Annex IX(I) Annex XI(A)	Excluded Class III Medical Devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	assurance Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regula	tion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			conformity verification		
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex IX(I) Annex IX(II)	Excluded Class III Medical Devices excepted those classified in Class III only as incorporating medicinal substances according to Directive 2001/83/EC and/or composed of substances or a combination of substances that are absorbed by o locally dispersed in the human body.

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Excluded Class III Medical Devices			
		 2. 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) 	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)	Excluded Class III Medical Devices Exclusion of 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 	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		human body via a body orifice or the dermal route 2. Non-active non-implantable devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on a quality	Annex XI(B) Annex IX(I) Annex IX(I)	Excluded Class III Medical Devices			
		- MDN 1214 General non-active non-implantable devices used in health care and other non-active non-implantable devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		MDS 1001 Devices incorporating medicinal substances			Excluded derived from human blood or human plasma			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including: aseptic processing,			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
					ethylene oxide gas sterilisation (EOG), moist heat sterilisation, radiation sterilisation (gamma-ray, x-ray, electron beam, beta-ray,).			
		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 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						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather,						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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					
		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					
NV MEDCERT GmbH ilatuspool 2 0355 HAMBURG sermany	0482	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- A. Active devices					
		- - 1. Active implantable devices - MDA 0101 Active implantable devices for stimulation/inhibition/monitoring	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)			
		-	Conformity assessment	Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Active implantable devices MDA 0103 Active implantable devices supporting or replacing organ functions 	based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(II)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation - 2. Active non-implantable devices for imaging, 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex IX(I) Annex IX(I)				
		monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			assurance						
		 - 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 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 0301 Active non-implantable devices utilising ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)					
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	assurance Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	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 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 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 0306 Active non-implantable devices for 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		extra-corporal circulation, administration or removal of substances and haemapheresis	based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	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 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical device	S
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices 	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 0312 Other active non-implantable surgical devices 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 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 	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 0315 Software	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 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	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on product quality assurance				
		 - 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 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 0318 Other active non-implantable devices 	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)			
		- B. Non-active devices					
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)			
		-	Conformity assessment	Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants 	based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(II)				
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials	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)				
		- - 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 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)				
		 - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		 - 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 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)				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)				
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	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)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			Conformity assessment based on product quality assurance						
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)					
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices	assurance Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				
		- MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	management system Conformity assessment based on assessment of technical documentation	Annex XI(A)				
			Conformity assessment based on product quality assurance					
		 - 2. 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) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - 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 	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)				
			Conformity assessment based on product quality assurance					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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 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)				
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			including: aseptic processing; ethylene oxide gas sterilisation (EOG); low temperature steam an formaldehyde sterilisation; moist heat sterilisation; radiation sterilisation (gamma, x-ray, electron beam); sterilisation with hydrogen peroxide; sterilisation with liquid chemical sterilising agens; thermic sterilisation with dr heat			
		MDS 1006 Reusable surgical instruments						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		MDS 1007 Devices incorporating or consisting of nanomaterial						
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		MDT 2006 Devices manufactured using chemical						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
MDC MEDICAL DEVICE CERTIFICATION GMBH Kriegerstrasse 6 70191 STUTTGART Germany	0483	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 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 MDT 2013 Devices which have undergone reprocessing I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE						
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)					
		 - 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 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)					
		 - 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 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)					
		-	Conformity assessment	Annex IX(I)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	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 0303 Active non-implantable devices utilising hyperthermia/hypothermia	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Except external pacemakers and heart defibrillators			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0305 Active non-implantable devices for stimulation or inhibition	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		- 3. Active non-implantable therapeutic devices and	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 0307 Active non-implantable respiratory devices 	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)	Except hyperbaric chambers			
		- 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0308 Active non-implantable devices for wound and skin care	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	assurance 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 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	excluding in vitro ferti¬lisation (IVF and assisted reproductive technologies (ART)

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	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 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	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)				

lame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		- 3. Active non-implantable therapeutic devices and	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)	
		- B. Non-active devices			
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)	
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials 	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)				
		- - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				
			based on product quality assurance					
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 2. 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) 	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)				
		- - 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	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)				
		-	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDN 1214 General non-active non-implantable devices used in health care and other non-active non-implantable devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)	
		MDS 1001 Devices incorporating medicinal substances			
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam and formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), sterilisation with hydrogen peroxide, thermic sterilisation with dry heat
		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 or mainly absorbed or			

ame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			
		MDS 1013 Class III custom-made implantable devices			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		MDT 2006 Devices manufactured using chemical			
		processing			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
SLG PRÜF UND ZERTIFIZIERUNGS GMBH	0494	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 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 MDT 2013 Devices which have undergone reprocessing			excluding reprocessing of single-use devices			
SLG PROF UND ZERTIFIZIERUNGS GMBH Burgstädter Strasse 20 09232 Hartmannsdorf Germany	0494	 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	restricted to X-ray diagnostics, scintigraphy			

LIST OF B Name and address of the notified bodies	NOTIFIED UNDER DIRECTIVE : Regulat Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	es Conditions
		assurance Conformity assessment based on product conformity verification		
	 - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
	 - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance Conformity assessment based on product conformity verification					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
			based on product conformity verification					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I)	excluding devices for external whole-body hyperthermia therapy and hyperthermic perfusion			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			assurance Conformity assessment based on product conformity verification		
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality		excluding devices for emergency medicine and anesthesia

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance Conformity assessment based on product conformity verification				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance Conformity assessment based on product conformity verification				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	restricted to products for minimally invasive surgery		
		 - 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 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	excluding prostheses		

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 o	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			assurance Conformity assessment based on product conformity verification		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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 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		restricted to central gas supply according to EN ISO 7396
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
			Conformity assessment based on product conformity verification				
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)					
		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			restricted to products included in the scope		
		MDS 1011 Devices in systems or procedure packs			restricted to products included in the scope		
		MDS 1012 Products without an intended medical purpose listed in Annex XVI to Regulation (EU) 2017/745			without Regulation (EU) 2017/745 Appendix XVI paragraph 1.; 2.; 3.; 4. restricted to products included in the scope		
		MDT 2001 Devices manufactured using metal processing					
		MDT 2002 Devices manufactured using plastic processing					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2010 Devices manufactured using electronic components including communication devices					
		MDT 2011 Devices which require packaging, including labelling			excluding sterile packaging		
		MDT 2012 Devices which require installation, refurbishment					
		MDT 2013 Devices which have undergone reprocessing			restricted to products that need to be reprocessed for use, excludin single-use devices		
urofins Electric & Electronics Finland Oy L 47 Kivimiehentie 4 I-02150 Espoo. inland	0537	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 	Conformity assessment based on a quality management system	Annex IX(I) Annex XI(A)			
		- MDA 0201 Active non-implantable imaging devices utilising ionizing radiation	Conformity assessment based on product quality assurance				
		- 2. Active non-implantable devices for imaging.	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation	based on product quality assurance					
		 - 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 system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices that directly contact central nervous system or central circulatory system, active therapeutic devices with an integrated or incorporated diagnostic function which significantly determines the patient management by the device (e.g. closed loop systems or automated external defibrillators), and devices that are intended for controlling, monitoring or directly influencing the performance of active implantable devices are excluded.			
		 - 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 system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices that directly contact central nervous system or central circulatory system, active therapeutic devices with an integrated or incorporated diagnostic function which significantly determines the patient management by the device (e.g. closed loop systems or automated external defibrillators), and devices that are intended for controlling, monitoring or directly influencing the performance of active implantable devices are excluded.			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex XI(A)	Devices that directly contact central nervous system or central circulatory system, active therapeutic devices with an integrated or incorporated diagnostic function which significantly determines the patient management by the device (e.g. closed loop systems or automated external defibrillators), and devices that are intended for controlling, monitoring or directly influencing the performance of active implantable devices are excluded.		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Heater-cooler units (blood warmers) are excluded.		
		 - 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 product quality assurance	Annex XI(A)	Devices that directly contact central nervous system or central circulatory system, therapeutic devices with an integrated or incorporated diagnostic function which significantly determines the patient management by the device (e.g. closed loop systems or automated external defibrillators), and devices that are intended for		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
					controlling, monitoring or directly influencing the performance of active implantable devices are excluded.		
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices that are intended for controlling, monitoring or directly influencing the performance of active implantable devices are excluded.		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices that directly contact central nervous system or central circulatory system are excluded.		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Active prostheses and exoskeletons are excluded.			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance		Software intended to provide information, which is used to take decisions having an impact that may cause death or an irreversible deterioration of a person's state of health, and therapeutic devices with an integrated or incorporated diagnostic function, which significantly determines the patient management by the device e.g. closed loop systems or automated external defibrillators, are excluded.			
		 - - 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 system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices for sterilization are excluded.			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex XI(A)				

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0318 Other active non-implantable devices	Conformity assessment based on product quality assurance		
		- B. Non-active devices			
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex XI(A)	Other devices except sutures, staples, screws, wedges, plates, wires, pins, clips and connectors are excluded.
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	
		 - 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 system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Devices for dialysis are excluded.
		 2. Non-active non-implantable devices MDN 1204 Non-active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	Contact lenses and intraocular lenses are excluded.
		 - - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	
		 - 2. Non-active non-implantable devices MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex XI(A)	

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex XI(A)	Devices (or their products of metabolism) that are systemically absorbed by the human body are excluded.
		 - 2. Non-active non-implantable devices - MDN 1214 General non-active non-implantable devices used in health care and other non-active 	Conformity assessment based on a quality management system Conformity assessment based on product quality assurance	Annex IX(I) Annex XI(A)	
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			Devices other than those intended to come into contact with intact skin only are excluded.
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			- Processes covered: aseptic processing, ethylene oxide gas sterilisation (EOG),moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam) - Processes excluded: low temperature steam and formaldehyde sterilisation

LIST OF B	SODIE	S NOTIFIED UNDER DIRECTIVE : Regulati	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		MDS 1006 Reusable surgical instruments			
		MDS 1007 Devices incorporating or consisting of nanomaterial			Devices presenting a high or medium potential for internal exposure are excluded.
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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			Devices intended for controlling, monitoring or directly influencing the performance of the active implantable are excluded.
		MDS 1010 Devices with a measuring function			
		MDS 1011 Devices in systems or procedure packs			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		MDT 2006 Devices manufactured using chemical			

LIST OF E		NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		processing 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			Devices manufactured using materials of human origin and devices other than intended to come into contact with intact skin only are excluded.
		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			
		MDT 2013 Devices which have undergone reprocessing			
CERTIQUALITY S.r.I. /ia G. Giardino, 4 20123 - MILANO taly	0546	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
		- A. Active devices			
		general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	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(II) Annex XI(A)	Excluded Class III Medical Device Except those classified in Class I only as incorporating medicinal substances, according to Directiv 2001/83/EC Excluding all devices depending on a source of electric energy

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	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)	Excluded Class III Medical Devices Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC Excluding all devices depending on a source of electrical energy
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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)	Excluded Class III Medical Devices Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC Excluding all devices depending on a source of electrical energy
		- - 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)	Excluded Class III Medical Devices
		- B. Non-active devices			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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(II) Annex XI(A)	Excluded Class III Medical Devices Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainly absorbed
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)	Excluded Class III Medical Devices Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainly absorbed
		- - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainly absorbed
		-	Conformity assessment	Annex IX(I)	Excluded Class III Medical Devices

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for administration, channelling and removal of substances, including devices for dialysis 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)	Except those classified in Class III only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainly absorbed
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)	
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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(II) Annex XI(A)	Excluded Class III Medical Device Except those classified in Class II only as incorporating medicinal substances, according to Directiv 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or main absorbed
		- - 2. Non-active non-implantable devices	Conformity assessment based on a quality	- ()	Excluded Class III Medical Device Except those classified in Class II

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDN 1207 Non-active non-implantable diagnostic devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainl absorbed
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)	Excluded Class III Medical Device
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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(II) Annex XI(A)	Excluded Class III Medical Device Except those classified in Class II only as incorporating medicinal substances, according to Directive 2001/83/EC and/or utilising biological active coatings and/or materials or being wholly or mainl absorbed
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Device

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		disinfecting, cleaning and rinsing	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 2. 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) 	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)	Exluding devices for in vitro fertilisation (IFV) 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 	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)	
		 - 2. Non-active non-implantable devices - MDN 1214 General non-active non-implantable devices used in health care and other non-active 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		non-implantable devices	based on assessment of technical documentation Conformity assessment based on product quality assurance		
		MDS 1001 Devices incorporating medicinal substances			Excluding plasma and blood derivatives
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilisation (EOG) - low temperature steam moist heat sterilisation - radiation sterilisation (gamma-ray, x-ray, electron beam)
		MDS 1006 Reusable surgical instruments			
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly or mainly absorbed or locally dispersed in the human body or are intended to undergo a chemical change in the body			
		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			
		MDT 2001 Devices manufactured using metal			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		processing			
		MDT 2002 Devices manufactured using plastic processing			
		MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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			
		MDT 2011 Devices which require packaging, including labelling			
		MDT 2012 Devices which require installation, refurbishment			
		MDT 2013 Devices which have undergone reprocessing			Limited to reusable devices whic have to undergone reprocessing
GS FIMKO OY akomotie 8 380 HELSINKI	0598	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
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LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb, excluding ultrasound devices			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb, excluding audiometers

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb, excluding blood warmers

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy)	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II)	Up to class IIb, limited to extracorporeal shockwave therapy of limbs and joints and shockwave HIFU
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb, excluding hyperbaric chambers
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb, excluding surgical devices

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Up to class IIb
		 - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II)	Up to class IIb, limited to hospital beds, physiotherapy equipment, rehabilitation, patient positioning and transport devices

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification		Up to class IIb, limited to autoclaves			
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on type-examination	Annex X Annex IX(I)	Up to class IIb			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(II) Annex XI(A) Annex XI(B)	
		B. Non-active devices - - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	Up to class IIb
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)	Up to class IIb

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)	Up to class IIb			
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)	Up to class IIb			
		- - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Up to class IIb			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Aseptic Processing, Ethylene Oxide gas sterilization, Low temperature steam and formaldehyde sterilization, Low temperature H2O2 sterilization, Moist heat sterilization, Radiation sterilization (gamma, x-ray, electron beam)
		MDS 1007 Devices incorporating or consisting of nanomaterial			Up to class IIb
		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			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		MDT 2001 Devices manufactured using metal			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		processing				
		MDT 2002 Devices manufactured using plastic processing				
		MDT 2003 Devices manufactured using non-metal mineral processing (e.g. glass, ceramics)				
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)				
		MDT 2006 Devices manufactured using chemical processing				
		MDT 2008 Devices manufactured in clean rooms and associated controlled environments				
		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				
Berlin Cert Prüf- und Zertifizierstelle für Medizinprodukte GmbH Dovestraße 6 10587 Berlin Germany	0633	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE				
		- A. Active devices				
		2. Active non-implantable devices for imaging,	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I)		
		monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices	bacad on a quality	Annex IX(II) Annex XI(A)		

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		utilising non-ionizing radiation	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)				
		 - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices 	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)		
		- 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		stimulation or inhibition	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)				
		 - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Without active devices intended to administer and/or remove medicinal products, body liquids or other substances to or from the body if this is done in a manner that is potentially hazardous, taking account of the nature of the substances involved, of the part of the body concerned and of the mode of application (Rule 12 Annex VIII Regulation (EU) 2017/745)			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory 	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical devic	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		nose and throat	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)	
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical 	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)			
		- MDA 0313 Active non-implantable prostneses, devices for rehabilitation and devices for patient positioning and transport	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)				
		- 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0317 Active non-implantable devices for	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		cleaning, disinfection and sterilisation	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		- B. Non-active devices					
		- - 2. Non-active non-implantable devices	Conformity assessment based on type-examination Conformity assessment	Annex X Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable	based on type-examination	Annex X Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		instruments	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(A) Annex XI(B)				
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		MDS 1004 Devices which are also machinery as define in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and o						

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		the Council (1)			
		MDS 1006 Reusable surgical instruments			
		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			
		MDT 2001 Devices manufactured using metal processing			
		MDT 2002 Devices manufactured using plastic processing			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2008 Devices manufactured in clean rooms and associated controlled environments			
		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			
STITUT PRO TESTOVÁNI A CERTIFIKACI, s. (INSTITUTE FOR TESTING AND	1023	I. CODES REFLECTING THE DESIGN AND			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices Name and address of the notified ID Responsible for the following products Responsible for the Annexes or Conditions								
bodies	U	/Horizontal technical competence	following procedures or modules	articles of the directives	Conditions			
ERTIFICATION) merged with ex-NB 1390 ida Tomase Bati 299 ouky, 76302 ZLIN czech Republic		INTENDED PURPOSE OF THE DEVICE						
		- A. Active devices						
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				
			Conformity assessment based on product quality assurance					
		- 2. Active non-implantable devices for imaging,	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)				
		- 2. Active non-implantable devices for imaging.	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	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 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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 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	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 	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)				
		general active non-implantable devices	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 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		-	assurance Conformity assessment	Annex IX(I)				
		- 3. Active non-implantable therapeutic devices and general active non-implantable devices	based on a quality management system	Annex IX(II) Annex XI(A)				
		- MDA 0312 Other active non-implantable surgical devices	Conformity assessment based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		- MDA 0315 Software	Conformity assessment based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				
		- MDA 0316 Medical gas supply systems and parts thereof	based on assessment of technical documentation Conformity assessment					
			based on product quality assurance					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 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)				
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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)				
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 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 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)	Excluding IUD, breast implants and non-absorbable dermal fillers based on methylmethacrylate nad silicones			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 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 system	Annex IX(I) Annex IX(II) Annex XI(A)				
		-	Conformity assessment	Annex IX(I)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1204 Non-active non-implantable devices for wound and skin care	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices 	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)				
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	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)				
		-	Conformity assessment	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDN 1208 Non-active non-implantable instruments	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)			
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)			
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices fo	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		disinfecting, cleaning and rinsing	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		 - 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 	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)				
		- - 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 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)				
		MDS 1001 Devices incorporating medicinal substances MDS 1004 Devices which are also machinery as defined	8					
		in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		the Council (1)					
		MDS 1005 Devices in sterile condition			Aseptic processing, ethylene oxide gas sterilisation (EOG), moist hea sterilisation, radiation sterilisation (gamma, x-ray, electron beam)		
		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 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			Excluding products intended to be totally or partially introduced into the human body through surgically invasive means for the purpose of modifying the anatomy or fixation of body parts. Excluding equipmer intended to be used to reduce, remove or destroy adipose tissue, such as equipment for liposuction,		
					lipolysis or lipoplasty.		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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			including only devices manufactured using processing of materials of microbial origin		
		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					
ENTE CERTIFICAZIONE MACCHINE SRL Via Ca' Bella, 243/A - loc. Castello di Serravalle 40053 Valsamoggia (BO)	1282	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
aly							
		- 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 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	.,	Excluding class III medical devices Limited to video endoscopes		
			Conformity assessment based on product quality assurance				
		 - 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 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)	Excluding class III medical devices		
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III medical device		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		general active non-implantable devices	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 class III medical devices			
			Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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)	Excluding class III medical devices			
		 - 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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III medical devices			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 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 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	.,	Excluding class III medical devices Limited to infusion pump
		- - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		Excluding class III medical devices Ecluding hyperbaric chambers
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	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)	Excluding class III medical devices
		-	Conformity assessment	Annex IX(I)	Excluding class III medical devices

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 	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(II)	Excluding class III medical devices Limited to devices for patient positioning and transport
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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)	Excluding class III medical device
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Excluding class III medical device

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		general active non-implantable devices - MDA 0316 Medical gas supply systems and parts thereof	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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)	Excluding class III medical devices Limited to moist heat sterilizers
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	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)	Excluding class III medical devices
		- B. Non-active devices	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Excluding class III medical devices

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices					
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)		
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)	Excluding class III medical devices	
		 - 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 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)	Excluding class III medical devices	
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III medical devices	

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		wound and skin care	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)	Excluding class III medical device		
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	Excluding class III medical device		
		 - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding class III medical device		

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices					
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
			based on assessment of technical documentation Conformity assessment based on product quality assurance			
		 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 	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)	Excluding class III medical devices	
		- - 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 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)	Excluding class III medical devices	
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)				
		MDS 1005 Devices in sterile condition			Limited to aseptic processing,	

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
					ethylene oxide gas sterilisation (EOG), moist heat sterilisation, radiation sterilisation		
		MDS 1006 Reusable surgical instruments					
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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					
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices					
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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 2010 Devices manufactured using electronic components including communication devices			
		MDT 2011 Devices which require packaging, including labelling			
		MDT 2012 Devices which require installation, refurbishment			
		MDT 2013 Devices which have undergone reprocessing			Limited to reusable devices which have to undergone reprocessing
SLOVENIAN INSTITUTE OF QUALITY AND METROLOGY - SIQ Mašera - Spasi#eva ulica 10 1000 LJUBLJANA Slovenia	1304	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 	based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives	5		
			based on product quality assurance Conformity assessment based on product conformity verification	Annex X			
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(I) Annex XI(A) Annex XI(B)			
		 - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance Conformity assessment based on product conformity verification		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	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 0305 Active non-implantable devices for stimulation or inhibition 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Annex X and XI(B) for lasers only. Other annexes with no limitations.		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)		
			based on product conformity verification			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	Conformity assessment based on a quality management system	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Annex X and XI(B) for lasers only. Other annexes with no limitations.	

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		-	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on a quality	Annex IX(I)	
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0315 Software	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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 0318 Other active non-implantable devices 	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)			
		- B. Non-active devices	Conformity assessment	Annex IX(I)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)	
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)	
		- - 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 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)	Nonabsorbable sutures only
		- - 2. Non-active non-implantable devices	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Devices for dialysis excluded

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		substances, including devices for dialysis	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)				
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	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)				
		- 2. Non-active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		orthopaedic and rehabilitation devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance				
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)			
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)			
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - 2. Non-active non-implantable devices - MDN 1213 Non-active non-implantable devices 	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		- - 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 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)	
		MDS 1001 Devices incorporating medicinal substances			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Aseptic processing, filtration, steam, EtO, irradiation included
		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 or mainly absorbed or locally dispersed in the human body or are intended to undergo a chemical change in the body			Biologically active coatings/materials excluded
		MDS 1009 Devices incorporating software/utilising			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		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						
		MDS 1013 Class III custom-made implantable devices			Orthopaedic implantable devices only			
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		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 2010 Devices manufactured using electronic components including communication devices						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		MDT 2011 Devices which require packaging, including labelling						
		MDT 2012 Devices which require installation, refurbishment						
		MDT 2013 Devices which have undergone reprocessing						
BUREAU VERITAS ITALIA S.P.A. Viale Monza, 347 20126 - MILANO (MI) Italy	1370	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 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				
		 - 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 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)				
			Conformity assessment	Annex IX(I)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices for monitoring and/or diagnosis 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	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)	Excluded Class III Medical Devices				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0305 Active non-implantable devices for stimulation or inhibition	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		- 3. Active non-implantable therapeutic devices and	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 0307 Active non-implantable respiratory devices 	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)	Excluding hyperbaric chamber			
		- 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- MDA 0311 Active non-implantable devices	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 inerapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		devices for rehabilitation and devices for patient positioning and transport	based on assessment of technical documentation Conformity assessment based on product quality assurance		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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)	Excluded Class III Medical Devices
		 - 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 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)	
		 B. Non-active devices - 1. Non-active implants and long term surgically invasive devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- MDN 1103 Non-active dental implants and dental materials	based on assessment of technical documentation Conformity assessment based on product quality assurance						
		- - 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 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)	Excluded Class III Medical Devices				
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)	Excluded Class III Medical Devices				
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)					

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)	
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)	
		 - - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		 - 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 	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilisation (EOG) - low temperature steam moist heat sterilisation - radiatic sterilisation (gamma-ray, x-ray, electron beam)
		MDS 1006 Reusable surgical instruments			
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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 2010 Devices manufactured using electronic components including communication devices			
		MDT 2011 Devices which require packaging, including labelling			
		MDT 2012 Devices which require installation, refurbishment			
		MDT 2013 Devices which have undergone reprocessing			Limited to reusable devices whic have to undergone reprocessing
OLSKIE CENTRUM BADAN I ERTYFIKACJI S.A. . Pu#awska 469 2-844 Warszawa	1434	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- A. Active devices						
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)				
		 - 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 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)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 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 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 0301 Active non-implantable devices utilising ionizing radiation	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 0302 Active non-implantable devices utilising non-ionizing radiation	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)					
		-	Conformity assessment	Annex IX(I)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	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 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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0306 Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemapheresis	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0309 Active non-implantable ophthalmologic devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices	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 0312 Other active non-implantable surgical 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		devices	based on assessment of technical documentation Conformity assessment based on product quality assurance		
		general active non-implantable devices	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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(II) Annex XI(A)	Excluding active non-implantable devices for in vitro fertilisation (IVF) and assisted reproductive technologies (ART)
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			technical documentation Conformity assessment based on product quality assurance					
		 - 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 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 0318 Other active non-implantable devices 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- B. Non-active devices						
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				
			based on product quality assurance					
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants 	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)				
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1103 Non-active dental implants and dental materials 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)				
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 2. Non-active non-implantable devices MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	assurance 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)				
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	assurance 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)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 2. 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) 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 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 	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)					
		- - 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 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)					
		MDS 1001 Devices incorporating medicinal substances MDS 1003 Devices manufactured utilising tissues or							

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), sterilisatio with liquid chemical sterilising agents			
		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 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						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		MDS 1013 Class III custom-made implantable devices					
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device					
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					
		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 materials of human origin		
		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					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
SGS Belgium NV Noorderlaan 87 BE-2030 Antwerpen Belgium	1639	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE						
Beigium		 - A. Active devices - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)	Excluding surgically invasive devices for transient/short term use utilising ionizing radiation			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)				
		- Z. ACTIVE NON-IMDIANTADIE DEVICES TOF IMADING.	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 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 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 0301 Active non-implantable devices utilising ionizing radiation 	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(II)	Excluding surgically invasive devices for transient/short term use utilising ionizing radiation			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	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 	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0306 Active non-implantable devices for	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				
		extra-corporal circulation, administration or removal of substances and haemapheresis	technical documentation Conformity assessment based on product quality assurance					
		general active non-implantable therapeutic devices and	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	assurance 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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	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 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)					
			assurance Conformity assessment	Annex IX(I)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0318 Other active non-implantable devices	Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		- B. Non-active devices					
		- - 1. Non-active implants and long term surgically invasive devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding heart valves		
		and neurovascular implants	Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			
			Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		1. Non-active implants and long term surgically	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)			
		invasive devices	management system	Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1103 Non-active dental implants and dental materials	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		- - 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 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)	- Excluding breast implants - Excluding implants and long term invasive devices utilising ionizing radiation			
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				
		 - - 2. Non-active non-implantable devices - MDN 1202 Non-active non-implantable devices for administration, channelling and removal of 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		substances, including devices for dialysis	based on assessment of technical documentation					
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)				
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices 	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)	
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	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)	
		 - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments 	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 2. 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) 	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)				
		- 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	management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding non-active devices for ingestion			
		- 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilization - moist heat sterilization - radiation sterilization (gamma, x-ray & electron beam) - gas plasma sterilization			
		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 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						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		purpose listed in Annex XVI to Regulation (EU) 2017/745						
		MDS 1013 Class III custom-made implantable devices						
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		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 devices manufactured using processing of materials of human origin			
		MDT 2010 Devices manufactured using electronic components including communication devices						
		MDT 2011 Devices which require packaging, including						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		labelling						
		MDT 2012 Devices which require installation, refurbishment			Excluding reprocessed single use devices			
Kiwa Dare B.V. Vijzelmolenlaan 7 NL-3447 GX Woerden Netherlands	1912	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	-	Conformity assessment based on type-examination	Annex X Annex IX(I)				
		Conformity assessment based on a quality management system	Annex IX(II) Annex XI(A)					
		Co bas	Conformity assessment based on assessment of technical documentation	Annex XI(B)				
			Conformity assessment based on product quality assurance					
			Conformity assessment based on product conformity verification					
		2. Active non-implantable devices for imaging,	Conformity assessment based on type-examination	Annex X Annex IX(I)				
		monitoring and/or diagnosis	Conformity assessment based on a quality management system	Annex IX(II) Annex XI(A)				
			Conformity assessment based on assessment of	Annex XI(B)				

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices for monitoring and/or diagnosis	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					
			Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	Conformity assessment	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 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	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification		Limited to devices for administration and 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I)	Excluding inhalation anaesthesia devices, lung ventilators and heart-lung machines

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		 - 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 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)			
		- - 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	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)			
		-	Conformity assessment	Annex X			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		general active non-implantable devices - MDA 0318 Other active non-implantable devices	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Limited to ethylene oxide gas sterilisation and radiation sterilisation
		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			Only for active devices
		MDS 1010 Devices with a measuring function MDS 1011 Devices in systems or procedure packs			Only devices in systems, procedure packs are excluded
		MDS 1012 Products without an intended medical			Only for active devices.

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
	purpose listed in Annex XVI to Regulation (EU) 2017/745						
	MDT 2001 Devices manufactured using metal processing						
	MDT 2002 Devices manufactured using plastic processing						
	MDT 2008 Devices manufactured in clean rooms and associated controlled environments						
	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						
1936	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 0201 Active non-implantable imaging devices utilising ionizing radiation 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices			
		/Horizontal technical competence purpose listed in Annex XVI to Regulation (EU) 2017/745 MDT 2001 Devices manufactured using metal processing MDT 2002 Devices manufactured using plastic processing MDT 2008 Devices manufactured in clean rooms and associated controlled environments 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 1936 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 0201 Active non-implantable imaging devices utilising ionizing radiation	/Horizontal technical competence following procedures or modules purpose listed in Annex XVI to Regulation (EU) 2017/745	/Horizontal technical competence following procedures or modules articles of the directives purpose listed in Annex XVI to Regulation (EU) 2017/745			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	Excluded, only for annex X and XI B: Transcutaneous partial pressure monitoring equipment, Electroencephalographs, Blood pressure indirect, automatic and periodic measuring equipment, Direct blood pressure monitoring equipment, Multiparametric patient monitors, Medical device for registration and analysis of data from single and multiple channel electrocardiograps, Screening thermographic device for verification of human fever, Electromechanic blood pressure measuring device, Clinical thermometers			
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II)	Excluded Class III Medical Devices Excluded, only for Annex X and XI B: Medical diagnostric Nuclear magnetic risonance device, Device for ultrasonic diagnosis and monitoring, Audiometers.			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			based on product conformity verification						
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	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)	Excluded Class III Medical Devic				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devic Excluded, only for Annex X and X B: Short wave therapy device, Ultrasonic therapy device, Microwave therapy device, Infant phototherapy equipment.				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	based on product conformity verification Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devic				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		-	based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment		Excluded, only for Annex X and XI			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0305 Active non-implantable devices for stimulation or inhibition 	based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	B: Neuromuscular stimulators.			
		 - 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 	Conformity assessment based on type-examination 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)	Excluded Class III Medical Devices Excluded, only for Annex X and XI B : Hemodialysis, Hemodiafiltration and haemofiltration device, Infusion pumps and control device, Peritoneal dialysis device.			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex IX(II)	Excluded Class III Medical Devices Excluding hyperbaric chamber Excluded, only for Annex X and XI B: Pulmunary ventilators, Systems for anesthesia, Iperbaric chambers, Device for the respiratory therapy of apnea during sliping, Spirometers.		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance				
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	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)	Excluded Class III Medical Devices		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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)	Excluded Class III Medical Devices		
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	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)	Excluded Class III Medical Devices		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or 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 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	Conformity assessment based on type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product	Annex IX(I) Annex IX(II)	Excluded Class III Medical Devices Excluded, only for Annex X and XI B: Lifting equipment used to transfer disable patients.			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0314 Active non-implantable devices for processing and preservation of human cells, tissues 	conformity verification Conformity assessment based on type-examination Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)	Excluded Class III Medical Devices Excluded in vitro fertilisation (IVF) and assisted reproductive technologies (ART)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART)	Conformity assessment based on assessment of technical documentation Conformity assessment					
			based on product quality assurance Conformity assessment based on product conformity verification					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices			
		 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0316 Medical gas supply systems and parts thereof 	assurance 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)	Excluded Class III Medical Devices			
		-	Conformity assessment	Annex IX(I)	Excluded Class III Medical Devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0318 Other active non-implantable devices	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)	Excluded Class III Medical Devices		
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)	Excluded neurovascular implant		
		-	Conformity assessment	Annex IX(I)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)	Excluded Class III Medical Devices			
		- - 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 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)				
		- - 2. Non-active non-implantable devices	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)	Excluded Class III Medical Devices			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)					
		 - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		Excluded dialysis devices in class				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)					
		- 2. Non-active non-implantable devices	Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices				

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	tion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		wound and skin care	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)	Excluded Class III Medical Devices
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)	Excluded Class III Medical Devices
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regula	tion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on assessment of technical documentation		
			Conformity assessment based on product quality assurance		
		2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments -	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on a quality	()	Excluded Class III Medical Devices
		 - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials 	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)	
		 - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices fo disinfecting, cleaning and rinsing 	Conformity assessment based on a quality management system r Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	Excluded Class III Medical Devices

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			technical documentation Conformity assessment based on product quality assurance				
		 - 2. 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) 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	()	Excluded Class III Medical Devices Excluded 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 	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)	Excluded Class III Medical Devices		
		- 2. Non-active non-implantable devices	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)	Excluded Class III Medical Devices		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on product quality assurance				
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)					
		MDS 1005 Devices in sterile condition			Including: - aseptic processing - ethylene oxide gas sterilisation (EOG) - low temperature steam - moist heat sterilisation - radiation sterilisation (gamma-ray, x-ray, electron beam)		
		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 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)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		2017/745					
		MDS 1013 Class III custom-made implantable devices			Limited to devices within the scope of designation related to the codes		
		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)					
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)					
		MDT 2005 Devices manufactured using biotechnology					
		MDT 2006 Devices manufactured using chemical processing					
		MDT 2008 Devices manufactured in clean rooms and associated controlled environments					
		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					
		MDT 2013 Devices which have undergone reprocessing			Limited to reusable devices which have to undergone reprocessing		
3EC International a.s. 3EC International a.s. Hranicna 18 Bratislava 82105 SLOVAKIA	2265	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical device	es
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
atislava 82105 ovakia					
		- A. Active devices			
		- 2. Active non-implantable devices for imaging, monitoring and/or diagnosis	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	
			technical documentation Conformity assessment based on product quality assurance		
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0203 Active non-implantable devices for monitoring of vital physiological parameters 	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)	
			Conformity assessment based on product quality		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices for monitoring and/or diagnosis 	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 		Annex IX(I) Annex IX(II) Annex XI(A)					
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	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 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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)				
			based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		general active non-implantable devices - MDA 0306 Active non-implantable devices for	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - 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 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 0308 Active non-implantable devices for wound and skin care 	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDA 0312 Other active non-implantable surgical devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance					
		 - 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 	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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 0315 Software 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0316 Medical gas supply systems and parts 	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 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	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			technical documentation Conformity assessment based on product quality assurance				
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)	excluding neurovascular implants		
		 - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants 	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)			
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		materials	technical documentation Conformity assessment based on product quality assurance		
		 - 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 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)	excluding breast implants
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)	
		 - 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices									
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			Conformity assessment based on product quality assurance						
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)					
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)					
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. 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)	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)				
		 - 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 	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)				
		- - 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 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)				
		MDS 1001 Devices incorporating medicinal substances						

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			excluding Reg. 722/2012
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam a formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam)
		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 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			except active implantable MDs
		MDS 1010 Devices with a measuring function			
		MDS 1011 Devices in systems or procedure packs			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		2017/745						
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		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						
		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						

Name and address of the notified bodies		S NOTIFIED UNDER DIRECTIVE : Regulat Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
TUV NORD Polska Sp. z o.o ul. Mickiewicza 29 40-085 Katowice Poland	2274	MDT 2013 Devices which have undergone reprocessing I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)	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 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)	
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding products used in ophthalmology

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		for monitoring and/or diagnosis	technical documentation Conformity assessment based on product quality assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	management system	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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)				
		- 3. Active non-implantable therapeutic devices and	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)	Including only infusion pumps, devices for dialysis, anaesthesia machines and devices for administration or removal of substances			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		 - 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 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)	Excluding hyperbaric chambers			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	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 0312 Other active non-implantable surgical devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 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 	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)	Excluding active prostheses			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software	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 0316 Medical gas supply systems and parts thereof 	Conformity assessment based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0317 Active non-implantable devices for cleaning, disinfection and sterilisation	assurance 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)				
		 B. Non-active devices 1. Non-active implants and long term surgically invasive devices MDN 1102 Non-active osteo- and orthopaedic implants 	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(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 on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		- - 1. Non-active implants and long term surgically invasive devices	management system	Annex IX(I) Annex IX(II) Annex XI(A)	Including only urological tapes, surgical meshes ligament and tendon prostheses made of			
		- MDN 1104 Non-active soft tissue and other implants	Conformity assessment based on assessment of technical documentation		multifilament polyester fibers			
			Conformity assessment based on product quality assurance					
		 - 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)				
			Conformity assessment based on product quality assurance					
		 - 2. Non-active non-implantable devices MDN 1204 Non-active non-implantable devices for wound and skin care 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)	Excluding sutures			
			Conformity assessment based on product quality assurance					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
		-	Conformity assessment	Annex IX(I)	Excluding devices for in vitro			

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		 - 2. 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) 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)	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 	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)	
		- - 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 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)	Including only ultrasound gels, medication cups
		MDS 1001 Devices incorporating medicinal substances			
		MDS 1005 Devices in sterile condition			Including: aseptic processing,

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
					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 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			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		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		
E Certiso Orvos- és Kórháztechnikai Illen#rz# és Tanúsító Kft. ird# u.101. iudakeszi lungary	2409	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
		- B. Non-active devices					
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)			

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			Conformity assessment based on product quality assurance						
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)					
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)					
		- 1. Non-active implants and long term surgically	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care 	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)				
		 - 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 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)				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care 	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)				
		 - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices 	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)				
		 - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices 	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)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)					
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)					
		- - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)					
		-	Conformity assessment	Annex IX(I)					

Name and address of the notified bodies ID Responsible for the following products (Horizontal technical competence) Responsible for the following procedures or articles of the directives Annexes or articles of the directives Conditions - 2. Non-active non-implantable devices - MIN 1211 Non-active non-implantable devices or a quality management system disinfecting, cleaning and rinsing Annex IX(II) Annex IX(II) Annex IX(II) - 2. Non-active non-implantable devices - MIN 1213 Non-active non-implantable devices Dased on a quality massesment based on assessment or active assessment based on assessment or active assessment based on assessment based on assessment or active assessment based on assessment or active assessment based on active assessment based on a quality management system Annex IX(II) - 2. Non-active non-implantable devices - MIN 1213 Non-active non-implantable devices Conformity assessment based on active transment based on a quality management system Annex IX(II) - 2. Non-active non-implantable devices - MIN 1213 Non-active non-implantable devices Conformity assessment based on a quality management system Annex IX(II) - 2. Non-active non-implantable devices - MIN 1214 General non-active non-implantable devices Annex IX(II) - 3. Non-active non-implantable devices MIN 1214 General non-active non-implantable Conformity assessment hased on a quality management system Annex IX(II)	LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical device	es
- MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing annex XI(A) Annex XI(A) - MDN 1211 Non-active non-implantable devices Conformity assessment based on product quality assurance Annex XI(A) - 2. Non-active non-implantable devices Conformity assessment based on a quality assessment based on sessessment of technical documentation Annex XI(A) - 2. Non-active non-implantable devices composed of substances to be introduced into the human body via a body orifice or the dermal route based on product quality assessment based on product quality assessment based on a quality management system Annex XI(A) - 2. Non-active non-implantable devices Conformity assessment based on product quality assessment based on product quality assessment based on product quality assessment based on a quality management system Annex XI(A) - 2. Non-active non-implantable devices Conformity assessment based on a quality assessment based on a routic quality assessment of rechnical documentation Annex XI(A) - 2. Non-active non-implantable devices Conformity assessment based on a quality assessment based on a routic quality assessment based on a routic quality assessment of rechnical documentation Annex XI(A) - A. Active devices Conformity assessment of rechnical documentation Annex XI(A)		ID		following procedures	articles of the	Conditions
- 2. Non-active non-implantable devices - ased on a quality Annex IX(II) - MDN 1213 Non-active non-implantable devices Conformity assessment Annex XI(A) - Composed of substances to be introduced into the human body via a body orifice or the dermal route Conformity assessment Annex XI(A) - Composed of substances to be introduced into the human body via a body orifice or the dermal route Conformity assessment Annex XI(A) - Composed of product quality assurance - Annex IX(II) Annex IX(II) - 2. Non-active non-implantable devices - Conformity assessment Annex IX(II) - 2. Non-active non-implantable devices - Annex IX(II) Annex IX(II) - 2. Non-active non-implantable devices - Conformity assessment Annex IX(II) - MDN 1214 General non-active non-implantable Conformity assessment Annex XI(A) - Onformity assessment Dased on a quality Annex XI(A) - MDN 1214 General non-active non-implantable Conformity assessment Annex XI(A) - Mon implantable devices - Conformity assessment Annex XI(A) - A. Active devices - A. Active devices - A. Active devices - A. Active devices			- MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality		
- 2. Non-active non-implantable devices based on a quality Annex IX(II) - MDN 1214 General non-active non-implantable Conformity assessment Annex XI(A) Conformity assessment based on assessment of Annex XI(A) Conformity assessment based on product quality Annex XI(A) - A. Active devices - A. Active devices Image: Active devices Image: Active devices			 - 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	
			 - 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	
Contormity accommont (Annov IV/I)				Conformity assessment	Annex IX(I)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - 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 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)					
		 - 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 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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0308 Active non-implantable devices for wound and skin care 	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 0310 Active non-implantable devices for ear, nose and throat 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	assurance Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 o	n medical devi	ces
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0311 Active non-implantable dental devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	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 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	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 0315 Software	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 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 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 0318 Other active non-implantable devices 	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B Name and address of the notified bodies	ID	S NOTIFIED UNDER DIRECTIVE : Regulat Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 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 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 0306 Active non-implantable devices for extra-corporal circulation, administration or removal of substances and haemapheresis 	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)	
		MDS 1001 Devices incorporating medicinal substances			excluding medicinal substances derived from human blood or human plasma
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG) low temperature steam ar formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), sterilisation with liquid chemical sterilising agents, sterilisation with hydrogen peroxide, sterilisation with dry he			
		MDS 1006 Reusable surgical instruments						
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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						
		MDS 1014 Devices incorporating as an integral part an						

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		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 human origin
		MDT 2010 Devices manufactured using electronic components including communication devices			
		MDT 2011 Devices which require packaging, including labelling			
NV Product Assurance AS eritasveien 1 363 Høvik lorway	2460	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
-		- A. Active devices			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	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)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)				
		 - 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 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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0204 Other active non-implantable devices for monitoring and/or diagnosis 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	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 0302 Active non-implantable devices utilising non-ionizing radiation 	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)					

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy) 	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 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 on product quality assurance	Annex IX(I) Annex IX(II) Annex XI(A)			
		3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		of substances and haemapheresis	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance				
		general active non-implantable devices - MDA 0307 Active non-implantable respiratory devices	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 0308 Active non-implantable devices for wound and skin care 	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	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		devices	based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		- - 3. Active non-implantable therapeutic devices and	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)			
		general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat	Conformity assessment based on assessment of technical documentation	Annex XI(A)			
			Conformity assessment based on product quality assurance				
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices	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 0312 Other active non-implantable surgical devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			technical documentation Conformity assessment based on product quality assurance				
		 - 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 	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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 0315 Software 	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)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on product quality assurance				
		 - 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 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 0318 Other active non-implantable devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on product quality assurance				
		- B. Non-active devices					
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	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)	Cardiac valves excluded		
			Conformity assessment based on product quality assurance				
		1. Non-active implants and long term surgically invasive devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			
		- MDN 1102 Non-active osteo- and orthopaedic implants	Conformity assessment based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		1. Non-active implants and long term surgically invasive devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			
		- MDN 1103 Non-active dental implants and dental materials	Conformity assessment based on assessment of technical documentation				
			Conformity assessment				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical devi	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			based on product quality assurance		
		 - 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 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)	
		- - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	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)	
		- - 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 system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance				
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)			
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic devices	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)				
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives				
		materials	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		- 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases	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)				
		- 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

- 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) Annex XI(A) - assisted reproductive technologies (ART) Conformity assessment based on product quality assurance Annex IX(I) - 2. Non-active non-implantable devices composed of subscripts of human body via a body onfice or the dermal route human body via a body onfice or the dermal route assurance Conformity assessment of echnical documentation conformity assessment based on a quality management system Annex IX(I) - 2. Non-active non-implantable devices composed of subscripts o	ame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
 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 based on a quality management system Conformity assessment of technical documentation Conformity assessment based on product quality assurance Annex IX(II) Annex XI(A) Annex IX(II) Annex XI(A) Annex XI(A) Annex IX(II) Annex XI(A) Annex XI(A) Conformity assessment based on a quality management system Conformity assessment of technical documentation Conformity assessment based on a quality management system Annex IX(II) 			processing and preservation of human cells, tissue or organs including in vitro fertilisation (IVF) and	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	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 - MDN 1214 General non-active non-implantable devices used in health care and other non-active non-implantable devices - MDN 1214 General non-active conformity assessment - MDN 1214 General non-active -			- MDN 1213 Non-active non-implantable devices composed of substances to be introduced into the	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	
assurance			- MDN 1214 General non-active non-implantable devices used in health care and other non-active	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(II)	

LIST OF B	BODIES	S NOTIFIED UNDER DIRECTIVE : Regulation	on (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives			
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)			
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam and formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), plasma sterilisation, chemical sterilisation and dry heat sterilisation
		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 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			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		MDS 1012 Products without an intended medical purpose listed in Annex XVI to Regulation (EU) 2017/745			All products without a medical purpose except: Devices intended to be totally or partially introduced into the human body through surgically invasive means for the purpose of fixation of body parts.			
		MDS 1013 Class III custom-made implantable devices						
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device						
		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)						
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)						
		MDT 2005 Devices manufactured using biotechnology						
		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						

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
JDEM Adriatic d.o.o.	2696	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 MDT 2013 Devices which have undergone reprocessing						
Radni#ka cesta 54/R3 Zagreb Croatia	2090	 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	()	EXCLUDING GAMMA RAY DEVICES			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			assurance				
		 - 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 system Conformity assessment based on product quality	Annex IX(I) 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 	assurance 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 0303 Active non-implantable devices utilising hyperthermia/hypothermia	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	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oı	n medical devid	ces
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy)	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance		
		 - 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 	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 0307 Active non-implantable respiratory devices 	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 0308 Active non-implantable devices for 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		wound and skin care	based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0309 Active non-implantable ophthalmologic devices 	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 0311 Active non-implantable dental devices 	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 0312 Other active non-implantable surgical devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical device	S
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 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 	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 0315 Software 	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 0316 Medical gas supply systems and parts thereof 	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)	

LIST OF B	BODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		 - 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 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 0318 Other active non-implantable devices 	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)	
		- B. Non-active devices			
		- 1. Non-active implants and long term surgically invasive devices	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 HEART VALVES

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 oi	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			Conformity assessment based on product quality assurance		
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)	
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)	
		 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 system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	EXCLUDING BREAST IMPLANTS

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care 	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)				
		 - 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 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)				
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions				
			assurance						
		- - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for wound and skin care	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)					
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)					
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	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)					

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical dev	ices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)	
		 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 	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)	
		-	Conformity assessment	Annex IX(I)	

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 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 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)			
		MDS 1001 Devices incorporating medicinal substances			EXCLUDING HUMAN BLOOD DERIVATIVES		
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)					
		MDS 1005 Devices in sterile condition			including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam and formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam)		
		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 or mainly absorbed or locally dispersed in the human body or are intended to undergo a chemical change in the body					

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			
		MDS 1013 Class III custom-made implantable devices			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		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			

LIST OF E	ODIES	NOTIFIED UNDER DIRECTIVE : Regulation	ion (EU) 2017/745 or	n medical dev	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			EXCLUDING REFURBISHMENT
		MDT 2013 Devices which have undergone reprocessing			
BSI Group The Netherlands B.V. Say Building, John M. Keynesplein 9, 1066 EP Amsterdam Netherlands	2797	I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE			
		- A. Active devices			
		 1. Active implantable devices MDA 0101 Active implantable devices for stimulation/inhibition/monitoring 	based on a quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		 1. Active implantable devices MDA 0102 Active implantable devices delivering drugs or other substances 	management system	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 1. Active implantable devices MDA 0103 Active implantable devices supporting or replacing organ functions 	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)				
		- - 1. Active implantable devices - MDA 0104 Active implantable devices utilising radiation and other active implantable devices	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)				
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices utilising ionizing radiation 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices utilising non-ionizing radiation 	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)				
		 - 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 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)				
		 - 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 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)				
		-	Conformity assessment	Annex IX(I)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0301 Active non-implantable devices utilising ionizing radiation 	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0302 Active non-implantable devices utilising non-ionizing radiation 	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 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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	Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF B	ODIES	NOTIFIED UNDER DIRECTIVE : Regulat	ion (EU) 2017/745 or	n medical devid	ces
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		general active non-implantable devices - MDA 0304 Active non-implantable devices for shock-wave therapy (lithotripsy)	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex X Annex IX(I) Annex IX(II) Annex XI(A) Annex XI(B)	
		 - - 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 	Conformity vernication Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			based on product quality assurance					
		 - - 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)				
			Conformity assessment based on product quality assurance					
		 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0308 Active non-implantable devices for wound and skin care 	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 0309 Active non-implantable ophthalmologic devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			assurance					
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0310 Active non-implantable devices for ear, nose and throat 	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 0311 Active non-implantable dental devices 	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 0312 Other active non-implantable surgical devices 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		 - 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 	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 0314 Active non-implantable devices for processing and preservation of human cells, tissues or organs including in vitrofertilisation (IVF) and assisted reproductive technologies (ART) 	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 0315 Software	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)				
		-	assurance Conformity assessment	Annex IX(I)				

LIST OF E	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		general active non-implantable devices	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	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 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 0318 Other active non-implantable devices	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)				
		- B. Non-active devices	Conformity assessment	Annex IX(I)				

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		- MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care	based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex IX(II) Annex XI(A)				
		 - 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 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)				
		- - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools	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)				
			Conformity assessment based on a quality	Annex IX(I) Annex IX(II)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		- MDN 1204 Non-active non-implantable devices for wound and skin care	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance	Annex XI(A)			
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	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)			
		- - 2. Non-active non-implantable devices - MDN 1206 Non-active non-implantable ophthalmologic devices	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)			
		- - 2. Non-active non-implantable devices - MDN 1207 Non-active non-implantable diagnostic	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF E	BODIES	NOTIFIED UNDER DIRECTIVE : Regula	ation (EU) 2017/745 or	n medical de	vices
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		devices	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality		
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	assurance Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	assurance 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)	
		 - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the 	Conformity assessment based on type-examination Conformity assessment based on a quality	Annex X Annex IX(I) Annex IX(II) Annex XI(A)	Annex X and XI(B) Limited to ma condoms

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices								
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
		transmission of sexually transmitted diseases	management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification	Annex XI(B)				
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	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)				
		 2. 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) 	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)				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		 - 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 	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)			
		- - 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 type-examination Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance Conformity assessment based on product conformity verification		Annex X and XI(B) limited to gloves		
		 1. Non-active implants and long term surgically invasive devices MDN 1101 Non-active cardiovascular, vascular and neurovascular implants 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

Name and address of the notified bodies	ID	NOTIFIED UNDER DIRECTIVE : Regulat Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		- - 1. Non-active implants and long term surgically invasive devices - MDN 1102 Non-active osteo- and orthopaedic implants	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)	
		- 1. Non-active implants and long term surgically	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)	
		 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF B	LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions			
			Conformity assessment based on product quality assurance					
		MDS 1001 Devices incorporating medicinal substances						
		MDS 1002 Devices manufactured utilising tissues or cells of human origin, or their derivatives						
		MDS 1003 Devices manufactured utilising tissues or cells of animal origin, or their derivatives						
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)						
		MDS 1005 Devices in sterile condition			Including aseptic processing, ethylene oxide gas sterilisation (EOG), low temperature steam an formaldehyde sterilisation, moist heat sterilisation, radiation sterilisation (gamma, x-ray, electron beam), sterilisation with hydrogen peroxide, sterilization with liquid chemical sterilizing agents, thermic sterilization with dry heat			
		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 or mainly absorbed or						

ame and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
		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			
		MDS 1013 Class III custom-made implantable devices			
		MDS 1014 Devices incorporating as an integral part an in vitro diagnostic device			
		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)			
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather, paper)			
		MDT 2005 Devices manufactured using biotechnology			
		MDT 2006 Devices manufactured using chemical			
		processing			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
		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 MDT 2010 Devices manufactured using electronic components including communication devices MDT 2011 Devices which require packaging, including labelling					
ntertek Medical Notified Body AB orshamnsgatan 43, Box 1103 SE-164 22 Kista	2862	MDT 2012 Devices which require installation, refurbishment MDT 2013 Devices which have undergone reprocessing I. CODES REFLECTING THE DESIGN AND INTENDED PURPOSE OF THE DEVICE					
Sweden		 - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0201 Active non-implantable imaging devices 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II)			
		- - 2. Active non-implantable devices for imaging, monitoring and/or diagnosis - MDA 0202 Active non-implantable imaging devices	based on assessment of technical documentation Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		 - 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 system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)	
		 - 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 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 0301 Active non-implantable devices utilising ionizing radiation 	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II)	
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance				
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0303 Active non-implantable devices utilising hyperthermia/hypothermia 	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 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 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	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		extra-corporal circulation, administration or removal of substances and haemapheresis	based on assessment of technical documentation Conformity assessment based on product quality assurance			
		 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0307 Active non-implantable respiratory devices 3. Active non-implantable therapeutic devices and general active non-implantable devices MDA 0308 Active non-implantable devices for wound and skin care 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation 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 IX(I) Annex IX(II) Annex XI(A)		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0311 Active non-implantable dental devices 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		-	assurance Conformity assessment based on a quality	Annex IX(I)		
		 - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0312 Other active non-implantable surgical devices 	management system Conformity assessment based on assessment of technical documentation	Annex IX(II) Annex XI(A)		
			Conformity assessment based on product quality assurance			
		- - 3. Active non-implantable therapeutic devices and general active non-implantable devices	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)		
		 MDA 0313 Active non-implantable prostheses, devices for rehabilitation and devices for patient positioning and transport 	Conformity assessment based on assessment of technical documentation			
			Conformity assessment based on product quality assurance			
		 - - 3. Active non-implantable therapeutic devices and general active non-implantable devices - MDA 0315 Software 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)		
			technical documentation Conformity assessment based on product quality			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices					
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or Conditions articles of the directives	
			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 0318 Other active non-implantable devices 	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)	
		- 1. Non-active implants and long term surgically	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)	
		- 1. Non-active implants and long term surgically	Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		- MDN 1102 Non-active osteo- and orthopaedic implants	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance			
		 1. Non-active implants and long term surgically invasive devices MDN 1103 Non-active dental implants and dental materials 	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)		
		 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 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)		
		 - 2. Non-active non-implantable devices - MDN 1201 Non-active non-implantable devices for anaesthesia, emergency and intensive care 	Conformity assessment based on a quality management system Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)		

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices							
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions		
			based on assessment of technical documentation				
			Conformity assessment based on product quality assurance				
		 - 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 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)			
		 - 2. Non-active non-implantable devices - MDN 1203 Non-active non-implantable guide catheters, balloon catheters, guidewires, introducers, filters, and related tools 	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)			
		 - 2. Non-active non-implantable devices - MDN 1204 Non-active non-implantable devices for 	Conformity assessment based on a quality management system Conformity assessment based on assessment of	Annex IX(I) Annex IX(II) Annex XI(A)			

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices					
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions
			technical documentation Conformity assessment based on product quality assurance		
		- - 2. Non-active non-implantable devices - MDN 1205 Non-active non-implantable orthopaedic and rehabilitation devices	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality	Annex IX(I) Annex IX(II) Annex XI(A)	
		- - 2. Non-active non-implantable devices - MDN 1208 Non-active non-implantable instruments	assurance 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)	
		- - 2. Non-active non-implantable devices - MDN 1209 Non-active non-implantable dental materials	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)	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
			Conformity assessment based on product quality assurance			
		 - 2. Non-active non-implantable devices - MDN 1210 Non-active non-implantable devices used for contraception or prevention of the transmission of sexually transmitted diseases 	Conformity assessment based on a quality management system Conformity assessment based on assessment of technical documentation	Annex IX(I) Annex IX(II)	Code scope limited to male condoms.	
		- - 2. Non-active non-implantable devices - MDN 1211 Non-active non-implantable devices for disinfecting, cleaning and rinsing	Conformity assessment based on assessment of technical documentation Conformity assessment	Annex IX(I) Annex IX(II) Annex XI(A)		
		- - 2. Non-active non-implantable devices - MDN 1213 Non-active non-implantable devices	based on product quality assurance Conformity assessment based on a quality management system	Annex IX(I) Annex IX(II) Annex XI(A)		
		composed of substances to be introduced into the human body via a body orifice or the dermal route	Conformity assessment based on assessment of technical documentation Conformity assessment based on product quality assurance			
		MDS 1001 Devices incorporating medicinal substances			Restricted to Article 117 devices.	

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		MDS 1004 Devices which are also machinery as defined in point (a) of the second paragraph of Article 2 of Directive 2006/42/EC of the European Parliament and of the Council (1)				
		MDS 1005 Devices in sterile condition			Including ethylene oxide gas sterilisation (EtO, EOG, moist heat sterilisation, aseptic processing, radiation sterilisation (gamma, x-ray, electron beam)	
		MDS 1006 Reusable surgical instruments				
		MDS 1008 Devices utilising biologically active coatings and/or materials or being wholly 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				
		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)				
		MDT 2004 Devices manufactured using non-metal non-mineral processing (e.g. textiles, rubber, leather,				

LIST OF BODIES NOTIFIED UNDER DIRECTIVE : Regulation (EU) 2017/745 on medical devices						
Name and address of the notified bodies	ID	Responsible for the following products /Horizontal technical competence	Responsible for the following procedures or modules	Annexes or articles of the directives	Conditions	
		paper)				
		MDT 2005 Devices manufactured using biotechnology				
		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				
		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				