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Approved Body: TUV SUD BABT Unlimited 0168

Legislation: Medical Devices Regulations 2002 (SI 2002 No 618, as amended) (UK

MDR 2002).

Reference: Medical Devices

The body is formally accredited against:

EN ISO/IEC 17021 - Certification of management systems

Name of National Accreditation Body (NAB): UKAS - United Kingdom

Accreditation Service

The accreditation covers the product categories and conformity assessment procedures concerned by this notification: Yes

Product family, product	Procedure/Modules	Annexes	Limitations
/Intended use/Product		or	
range		articles of	
_		the	
		directives	
		[as	
		modified	
		by	
		Part III of	

		Schedule 2A to the Medical Devices Regulations 2002]	
MD 0100 General non- active, non-implantable medical devices			
MD 0101 – Non-active devices for anaesthesia, emergency and intensive care	Full quality assurance system; Production quality assurance; Product quality assurance	Annex II; Annex V; Annex VI	
MD 0102 – Non-active devices for injection, infusion, transfusion and dialysis	Full quality assurance system; Production quality assurance; Product quality	Annex II; Annex V; Annex VI	
MD 0103 – Non-active orthopaedic and rehabilitation devices	assurance Full quality assurance system; Production quality assurance; Product quality assurance	Annex II; Annex V; Annex VI	
MD 0104 – Non-active medical devices with measuring function	Full quality assurance system; Production quality assurance; Product quality assurance	Annex II; Annex V; Annex VI	
MD 0105 – Non-active ophthalmologic devices	Full quality assurance system; Production quality assurance; Product quality assurance	Annex II; Annex V; Annex VI	
MD 0106 – Non-active instruments	Full quality assurance system; Production quality assurance;	Annex II; Annex V;	

	Product quality assurance	Annex VI
MD 0107 – Contraceptive medical devices	Full quality assurance system; Production quality	Annex II; Annex V;
	assurance; Product quality assurance	Annex VI
MD 0108 – Non-active medical devices for	Full quality assurance	Annex II;
disinfecting, cleaning, rinsing	system; Production quality assurance;	Annex V;
	Product quality assurance	Annex VI
MD 0109 – Non-active devices for in vitro	Full quality assurance system;	Annex II;
fertilisation (IVF) and assisted reproductive	Production quality assurance;	Annex V;
technologies (ART)	Product quality assurance	Annex VI
MD 0110 – Non-active medical devices for ingestion	Full quality assurance system;	Annex II;
3	Production quality assurance;	Annex V;
	Product quality assurance	Annex VI
MD 0200 - Non-active implants		
MD 0201 – Non-active cardiovascular implants	Full quality assurance system;	Annex II;
·	Production quality assurance;	Annex V;
	Product quality assurance	Annex VI
MD 0202 – Non-active orthopaedic implants	Full quality assurance system;	Annex II;
	Production quality assurance;	Annex V;
	Product quality assurance	Annex VI
MD 0203 – Non-active functional implants	Full quality assurance system;	Annex II;
·	Production quality assurance;	Annex V;
	Product quality assurance	Annex VI

MD 0204 – Non-active soft	Full quality assurance	Annex II;	Excluding breast
tissue implants	system; Production quality	Annex V;	implants.
	assurance; Product quality assurance	Annex VI	
MD 0300 – Devices for wound care			
MD 0301 – Bandages and wound dressings	Full quality assurance system;	Annex II;	
	Production quality assurance; Product quality	Annex V; Annex VI	
	assurance	-	
MD 0302 – Suture material and clamps	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 0303 – Other medical devices for wound care	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 0400 – Non-active dental devices and			
accessories			
MD 0401 – Non-active dental equipment and instruments	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 0402 – Dental materials	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 0403 – Dental implants	Full quality assurance	Annex II;	
	system; Production quality	Annex V;	
	assurance; Product quality	Annex VI	
	assurance		

MD 1100 – General active medical devices			
MD 1101 – Devices for extra- corporal circulation, infusion	Full quality assurance system;	Annex II;	
and haemopheresis	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1102 – Respiratory devices, devices including	Full quality assurance system;	Annex II;	
hyperbaric chambers for oxygen therapy, inhalation	Production quality assurance;	Annex V;	
anaesthesia	Product quality assurance	Annex VI	
MD 1103 – Devices for stimulation or inhibition	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex VI	
MD 4404 Astive averical	Product quality assurance	-	
MD 1104 – Active surgical devices	Full quality assurance system; Production quality	Annex II; Annex V;	
	assurance; Product quality	Annex VI	
MD 1105 – Active	assurance Full quality assurance	Annex II;	
ophthalmologic devices	system; Production quality	Annex V;	
	assurance; Product quality assurance	Annex VI	
MD 1106 – Active dental devices	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1107 – Active devices for disinfection and sterilisation	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1108 – Active rehabilitation devices and	Full quality assurance system;	Annex II;	
active prostheses	Production quality assurance;	Annex V;	

	Product quality assurance	Annex VI	
MD 1109 – Active devices for patient positioning and transport	Full quality assurance system; Production quality	Annex II; Annex V;	
	assurance; Product quality assurance	Annex VI	
MD 1110 – Active devices for in vitro fertilisation (IVF) and assisted reproductive	Full quality assurance system; Production quality	Annex II; Annex V;	
technologies (ART)	assurance; Product quality assurance	Annex VI	
MD 1111 – Software	Full quality assurance system; Production quality	Annex II; Annex V;	
	assurance; Product quality assurance	Annex VI	
MD 1112 – Parts incorporated within Medical gas supply systems	Full quality assurance system; Production quality	Annex II; Annex V;	Excluding medical gas pipelines (Not considered to be
	assurance; Product quality assurance	Annex VI	medical devices). Including parts of the
			system such as regulators and valves.
MD 1200 – Devices for imaging			
MD 1201 – Imaging devices utilising ionizing radiation	Full quality assurance system;	Annex II;	
	Production quality assurance; Product quality assurance	Annex V; Annex VI	
MD 1202 – Imaging devices utilising non-ionizing	Full quality assurance system;	Annex II;	
radiation	Production quality assurance; Product quality	Annex V; Annex VI	
MD 1300 – Monitoring devices	assurance		
ucvices			

MD 1301 – Monitoring devices of non-vital	Full quality assurance system;	Annex II;	
physiological parameters	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1302 – Monitoring devices of vital physiological	Full quality assurance system;	Annex II;	
parameters	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1400 – Devices for radiation therapy and thermotherapy			
MD 1401 – Devices utilising ionizing radiation	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1402 – Devices utilising non-ionizing radiation	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1403 – Devices for hyperthermia	Full quality assurance system;	Annex II;	
	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
MD 1404 – Devices for (extracorporal) shock-wave	Full quality assurance system;	Annex II;	
therapy (lithotripsy)	Production quality assurance;	Annex V;	
	Product quality assurance	Annex VI	
			Limitationo
Horizontal technical competence			Limitations
MDS 7001 – Medical devices incorporating medicinal substances, according to the Human Medicines Regulations 2012			
MDS 7002 – Medical devices utilising tissues of animal origin, including Commission Regulation (EU) No 722/2012			
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MDS 7003 – Medical devices incorporating derivates of human blood, according to UK Medical Devices Regulations 2002	
MDS 7004 – Medical devices referencing The Supply of Machinery (Safety) Regulations 2008	
MDS 7006 – Medical devices in sterile condition	
MDS 7008 – Medical devices utilising nanomaterials	
MDS 7009 – Medical devices utilising biological active coating and/or materials or being wholly or mainly absorbed	
MDS 7010 – Medical devices incorporating software / utilising software / controlled by software	