MDS - G020

قائمة المواصفات القياسية المعترف بها (لدعم طلبات الإذن بتسويق الأجهزة و المستلزمات الطبية)

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صفحة 1 من103

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مقدمة

تهدف قائمة المواصفات القياسية المعترف بها مساعدة مقدمي طلبات الإذن بتسويق الأجهزة و المستلزمات الطبية على إستيفاء المتطلبات اللازمة للحصول على أذن بتسويق الأجهزة والمستلزمات الطبية وذلك عن طريق إثبات إمتثال الجهاز أو المستلزم الطبي للمبادئ الأساسية للسلامة والأداء وذلك خلال دورة حياة الجهاز الطبي كاملة. وقد تم تصنيف قوائم المواصفات بناء على مجال عمل الأجهزة والمستزلمات الطبية وحصر المواصفات العامة والتي تنطبق على اغلب الأجهزة والمتسلزمات الطبية في مجال واحد.

طريقة شراء المواصفة

يمكن الحصول على المواصفات القياسية السعودية المعتمدة من الهيئة العامة للغذاء والدواء وذلك بنسخ رقمية عن طريق المتحر الإلكتروني للمواصفة القياسية يمكن زيارة الموقع الخاص بالمنظمة الدولية المصدرة للمواصفة.

- رابط متجر المنظمة الدولية لتقييس (ISO)
- رابط متجر المنظمة الدولية الكهروتقنية (IEC)

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10.	ISO 9360-2:2001 Anaesthetic and respiratory equipment - Heat and moisture exchangers (HMEs) for humidifying respired gases in humans - Part 2: HMEs for use with tracheostomized patients having minimum tidal volumes of 250 ml
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28.	ISO 81060-1:2007 Non-invasive sphygmomanometers - Part 1: Requirements and test methods for non-automated measurement type

29.	ISO 80601-2-55:2018 Medical electrical equipment – Part 2-55: Particular requirements for the basic safety and essential performance of respiratory gas monitors
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32.	ISO 5360:2016 Anaesthetic vaporizers — Agent-specific filling systems
33.	ISO 27427:2023 Anaesthetic and respiratory equipment Nebulizing systems and components
34.	ISO 18250-1:2018 Medical devices — Connectors for reservoir delivery systems for healthcare applications — Part 1: General requirements and common test methods
35.	IEC 60601-1-6:2010+AMD1:2013+AMD2:2020 Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability
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58.	ISO 21917:2021 Anaesthetic and respiratory equipment — Voice prostheses
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60.	ISO 23371:2022 Anaesthetic and respiratory equipment — Cuff pressure indication, control and regulation devices
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62.	ISO 8835-7:2011 Inhalational anaesthesia systems — Part 7: Anaesthetic systems for use in areas with limited logistical supplies of electricity and anaesthetic gases
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64.	ISO 18835:2015 Inhalational anaesthesia systems — Draw-over anaesthetic systems
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95.	SASO-IEC-60086-4:2007 Primary batteries - Part 4: Safety of lithium batteries"

96.	SASO-IEC-62281:2018 Safety of primary and secondary lithium cells and batteries during transport
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6.	ISO/TS 23128:2019 Medical devices — Transfusion set and blood bag compatibility test method
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	ISO 8536-9:2015
50.	Infusion equipment for medical use — Part 9: Fluid lines for single use with pressure infusion equipment
	ISO 8536-8:2015
51.	Infusion equipment for medical use — Part 8: Infusion sets for single use with pressure infusion apparatus
	ISO 8536-7:2009
52.	Infusion equipment for medical use — Part 7: Caps made of aluminium-plastics combinations for infusion bottles
	ISO 8536-6:2016
53.	Infusion equipment for medical use — Part 6: Freeze drying closures for infusion bottles
<i>51</i>	ISO 8536-5:2004
54.	Infusion equipment for medical use — Part 5: Burette infusion sets for single use, gravity feed
5.5	ISO 8536-4:2019 5 F D A
55.	Infusion equipment for medical use — Part 4: Infusion sets for single use, gravity feed
	ISO 8536-3:2009
56.	Infusion equipment for medical use — Part 3: Aluminium caps for infusion bottles
	ISO 8536-2:2023
57.	Infusion equipment for medical use — Part 2: Closures for infusion bottles
50	ISO 8536-1:2011
58.	Infusion equipment for medical use — Part 1: Infusion glass bottles
	ISO 8362-7:2006
59.	Injection containers and accessories — Part 7: Injection caps made of aluminium-plastics combinations without overlapping plastics part

	ISO 8362-6:2010
60.	Injection containers and accessories — Part 6: Caps made of aluminium-plastics combinations for injection vials
	ISO 8362-5:2016
61.	Injection containers and accessories — Part 5: Freeze drying closures for injection vials
	ISO 8362-4:2011
62.	Injection containers and accessories — Part 4: Injection vials made of moulded glass
	ISO 8362-3:2001
63.	Injection containers and accessories — Part 3: Aluminium caps for injection vials
	ISO 8362-2:2015
64.	Injection containers and accessories — Part 2: Closures for injection vials
	ISO 8362-1:2018
65.	Injection containers and accessories — Part 1: Injection vials made of glass tubing
	ISO 6710:2017
66.	Single-use containers for human venous blood specimen collection
	ISO 4802-2:2016
67.	Glassware — Hydrolytic resistance of the interior surfaces of glass containers — Part 2: Determination by flame spectrometry and classification
	ISO 4802-1:2016
68.	Glassware — Hydrolytic resistance of the interior surfaces of glass containers — Part 1:
	Determination by titration method and classification
	ISO 3826-4:2015
69.	Plastics collapsible containers for human blood and blood components — Part 4: Aphaeresis blood bag systems with integrated features
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	ISO 3826-3:2006
70.	Plastics collapsible containers for human blood and blood components — Part 3: Blood bag systems with integrated features
	ISO 3826-2:2008
71.	Plastics collapsible containers for human blood and blood components — Part 2: Graphical symbols for use on labels and instruction leaflets
	ISO 3826-1:2019
72.	Plastics collapsible containers for human blood and blood components — Part 1: Conventional containers
	ISO 3749:2022
73.	Glass syringes — Determination of extractable tungsten
	ISO 1135-5:2015
74.	Transfusion equipment for medical use — Part 5: Transfusion sets for single use with pressure infusion apparatus
	ISO 1135-4:2015
75.	Transfusion equipment for medical use — Part 4: Transfusion sets for single use, gravity feed
	ISO 1135-3:2016
76.	Transfusion equipment for medical use — Part 3: Blood-taking sets for single use
	ISO 720:2020
77.	Glass — Hydrolytic resistance of glass grains at 121 °C — Method of test and classification
	ISO 719:2020
78.	Glass — Hydrolytic resistance of glass grains at 98 °C — Method of test and classification

	Biological Evaluation
1.	ISO/TR 10993-22:2017 Biological evaluation of medical devices — Part 22: Guidance on nanomaterials
2.	ISO 10993-23:2021 Biological evaluation of medical devices — Part 23: Tests for irritation
3.	ISO/TR 10993-33:2015 Biological evaluation of medical devices — Part 33: Guidance on tests to evaluate genotoxicity — Supplement to ISO 10993-3
4.	ISO/TR 10993-55:2023 Biological evaluation of medical devices — Part 55: Interlaboratory study on cytotoxicity
5.	ISO/TS 11796:2023 Biological evaluation of medical devices — Requirements for interlaboratory studies to demonstrate the applicability of validated in vitro methods to assess the skin sensitization of medical devices
6.	ISO/TR 21582:2021 Pyrogenicity — Principles and methods for pyrogen testing of medical devices
7.	ISO/TS 21726:2019 Biological evaluation of medical devices — Application of the threshold of toxicological concern (TTC) for assessing biocompatibility of medical device constituents
8.	ISO/TR 22442-4:2010 Medical devices utilizing animal tissues and their derivatives — Part 4: Principles for elimination and/or inactivation of transmissible spongiform encephalopathy (TSE) agents and validation assays for those processes

	ISO/TS 37137-1:2021
9.	Biological evaluation of absorbable medical devices — Part 1: General requirements
	ISO/TR 37137:2014
10.	Cardiovascular biological evaluation of medical devices — Guidance for absorbable implants



	Implantable Devices	
1.	ISO 7197:2006 + Cor 1:2007 Neurosurgical implants — Sterile, single-use hydrocephalus shunts and components	
2.	ISO 9713:2022 Neurosurgical implants — Self-closing intracranial aneurysm clips	
3.	ISO 13179-1:2021 Implants for surgery — Coatings on metallic surgical implants — Part 1: Plasma-sprayed coatings derived from titanium or titanium-6 aluminum-4 vanadium alloy powders	
4.	ISO/TR 14283:2018 Implants for surgery — Essential principles of safety and performance	
5.	ISO 14607:2018 Non-active surgical implants — Mammary implants — Particular requirements	
6.	ISO 14630:2012 Non-active surgical implants — General requirements	
7.	ISO 16054:2019 Implants for surgery — Minimum data sets for surgical implants	
8.	ISO 16061:2021 Instruments for use in association with non-active surgical implants — General requirements	
9.	ISO 17327-1:2018 Non-active surgical implants — Implant coating — Part 1: General requirements	
10.	ISO 19213:2017 Implants for surgery — Test methods of material for use as a cortical bone model	

	ISO 19227:2018
11.	Implants for surgery — Cleanliness of orthopedic implants — General requirements
	ISO 14708-1:2014
12.	Implants for surgery — Active implantable medical devices — Part 1: General requirements for safety, marking and for information to be provided by the manufacturer
	ISO 14708-2:2019
13.	Implants for surgery — Active implantable medical devices — Part 2: Cardiac pacemakers
	ISO 14708-3:2017
14.	Implants for surgery — Active implantable medical devices — Part 3: Implantable neurostimulators
	ISO 14708-4:2022
15.	Implants for surgery — Active implantable medical devices — Part 4: Implantable infusion pump systems
	ISO 14708-5:2020
16.	Implants for surgery — Active implantable medical devices — Part 5: Circulatory support devices
	ISO 14708-6:2019
17.	Implants for surgery — Active implantable medical devices — Part 6: Particular requirements for active implantable medical devices intended to treat tachyarrhythmia (including implantable defibrillators)
	ISO 14708-7:2019
18.	Implants for surgery — Active implantable medical devices — Part 7: Particular requirements for cochlear and auditory brainstem implant systems
	ISO 27185:2012
19.	Cardiac rhythm management devices — Symbols to be used with cardiac rhythm management device labels, and information to be supplied — General requirements
20.	IEC 60601-2-31:2020

	Medical electrical equipment — Part 2-31: Particular requirements for the basic safety and essential performance of external cardiac pacemakers with internal power source
21.	ISO 5840-1:2021 Cardiovascular implants - Cardiovascular implants - Cardiac valve prostheses - Part 1: General requirements
22.	ISO 5840-2:2021 Cardiovascular implants — Cardiac valve prostheses — Part 2: Surgically implanted heart valve substitutes
23.	ISO 5840-3:2021 Cardiovascular implants — Cardiac valve prostheses — Part 3: Heart valve substitutes implanted by transcatheter techniques
24.	ISO 7198:2016 Cardiovascular implants and extracorporeal systems — Vascular prostheses — Tubular vascular grafts and vascular patches
25.	ISO 7199:2016 + Amd 1:2020 Cardiovascular implants and artificial organs — Blood-gas exchangers (oxygenators)
26.	ISO 8637-1:2017 Extracorporeal systems for blood purification — Part 1: Haemodialysers, haemodiafilters, haemofilters and haemoconcentrators
27.	ISO 8637-2:2018 Extracorporeal systems for blood purification — Part 2: Extracorporeal blood circuit for haemodialysers, haemodiafilters and haemofilters
28.	ISO 8637-3:2018 Extracorporeal systems for blood purification — Part 3: Plasmafilters

	ISO 12417-1:2015
29.	Cardiovascular implants and extracorporeal systems — Vascular device-drug combination products — Part 1: General requirements
	ISO 15675:2016
30.	Cardiovascular implants and artificial organs — Cardiopulmonary bypass systems — Arterial blood line filters
	ISO 15676:2016
31.	Cardiovascular implants and artificial organs — Requirements for single-use tubing packs for cardiopulmonary bypass and extracorporeal membrane oxygenation (ECMO)
	ISO/TS 17137:2021
32.	Cardiovascular implants and extracorporeal systems — Cardiovascular absorbable implants
	ISO 18193:2021
33.	Cardiovascular implants and artificial organs — Cannulae for extracorporeal circulation
	ISO 23500-1:2019
34.	Preparation and quality management of fluids for haemodialysis and related therapies — Part 1: General requirements
	ISO 23500-2:2019
35.	Preparation and quality management of fluids for haemodialysis and related therapies — Part 2: Water treatment equipment for haemodialysis applications and related therapies
	ISO 23500-3:2019
36.	Preparation and quality management of fluids for haemodialysis and related therapies — Part 3: Water for haemodialysis and related therapies
	ISO 23500-4:2019
37.	Preparation and quality management of fluids for haemodialysis and related therapies — Part 4: Concentrates for haemodialysis and related therapies

	ISO 22500 5-2010
38.	ISO 23500-5:2019 Preparation and quality management of fluids for haemodialysis and related therapies — Part 5: Quality of dialysis fluid for haemodialysis and related therapies
39.	ISO 7206-1:2008 Implants for surgery — Partial and total hip joint prostheses — Part 1: Classification and designation of dimensions
40.	ISO 7206-2:2011+ Amd 1:2016 Implants for surgery — Partial and total hip joint prostheses — Part 2: Articulating surfaces made of metallic, ceramic and plastics materials
41.	ISO 7206-6:2013 Implants for surgery — Partial and total hip joint prostheses — Part 6: Endurance properties testing and performance requirements of neck region of stemmed femoral components
42.	ISO 7206-10:2018+ Amd 1:2021 Implants for surgery — Partial and total hip-joint prostheses — Part 10: Determination of resistance to static load of modular femoral heads
43.	ISO 7206-12:2016 Implants for surgery — Partial and total hip joint prostheses — Part 12: Deformation test method for acetabular shells
44.	ISO 7206-13:2016+ Amd 1:2022Implants for surgery — Partial and total hip joint prostheses — Part 13: Determination of resistance to torque of head fixation of stemmed femoral components
45.	ISO 21534:2007 Non-active surgical implants — Joint replacement implants — Particular requirements
46.	ISO/TS 20721:2020 Implants for surgery — General guidelines and requirements for assessment of absorbable metallic implants

47.	ISO 22926:2023 Implants for surgery — Specification and verification of synthetic anatomical bone models for testing
	implants for surgery — specification and verification of synthetic anatomical bone models for testing
48.	ISO 5832-1:2016
	Implants for surgery — Metallic materials — Part 1: Wrought stainless steel
	ISO 5832-2:2018
49.	Implants for surgery — Metallic materials — Part 2: Unalloyed titanium
	ISO 5832-3:2021
50.	Implants for surgery — Metallic materials — Part 3: Wrought titanium 6-aluminium 4-vanadium alloy
	ISO 5832-4:2014
51.	Implants for surgery — Metallic materials — Part 4: Cobalt-chromium-molybdenum casting alloy
	ISO 5832-5:2022
52.	Implants for surgery — Metallic materials — Part 5: Wrought cobalt-chromium-tungsten-nickel
	ISO 5832-6:2022 5 F D A
53.	Implants for surgery — Metallic materials — Part 6: Wrought cobalt-nickel-chromium-molybdenum alloy
	ISO 5832-7:2016
54.	Implants for surgery — Metallic materials — Part 7: Forgeable and cold-formed cobalt-chromium-nickel-molybdenum-iron alloy
	ISO 5832-9:2019
55.	Implants for surgery — Metallic materials — Part 9: Wrought high nitrogen stainless steel
	ISO 5832-11:2014
56.	Implants for surgery — Metallic materials — Part 11: Wrought titanium 6-aluminium 7-niobium alloy

	ISO 5832-12:2019
57.	Implants for surgery — Metallic materials — Part 12: Wrought cobalt-chromium-molybdenum alloy
58.	ISO 5832-14:2019 Implants for surgery — Metallic materials — Part 14: Wrought titanium 15-molybdenum 5-zirconium 3-aluminium alloy
59.	ISO 5833:2002 Implants for surgery — Acrylic resin cements
60.	ISO 6474-1:2019 Implants for surgery — Ceramic materials — Part 1: Ceramic materials based on high purity alumina
61.	ISO 6474-2:2019 Implants for surgery — Ceramic materials — Part 2: Composite materials based on a high-purity alumina matrix with zirconia reinforcement
62.	ISO 13779-4:2018 Implants for surgery — Hydroxyapatite — Part 4: Determination of coating adhesion strength
63.	ISO 13782:2019 Implants for surgery — Metallic materials — Unalloyed tantalum for surgical implant applications
64.	ISO 15374:1998 Implants for surgery — Requirements for production of forgings
65.	ISO 16402:2008 Implants for surgery — Acrylic resin cement — Flexural fatigue testing of acrylic resin cements used in orthopaedics
66.	ISO/TS 21560:2020 General requirements of tissue-engineered medical products

Prosthetics and Orthotics	
1.	ISO 10328:2016 Prosthetics - Structural testing of lower-limb prostheses - Requirements and test methods
2.	ISO 22523:2006 External limb prostheses and external orthoses - Requirements and test methods
3.	ISO 22675:2016 Prosthetics - Testing of ankle-foot devices and foot units - Requirements and test methods
4.	ISO/TS 4549:2023 Orthotics -Method for testing the reliability of microprocessor-controlled ankle moment units of ankle-foot orthoses
5.	ISO 8548-1:1989 Prosthetics and orthotics — Limb deficiencies — Part 1: Method of describing limb deficiencies present at birth
6.	ISO 8548-2:2020 Prosthetics and orthotics — Limb deficiencies — Part 2: Method of describing lower limb amputation stumps
7.	ISO 8548-3:1993 Prosthetics and orthotics — Limb deficiencies — Part 3: Method of describing upper limb amputation stumps
8.	ISO 8548-4:1998 Prosthetics and orthotics — Limb deficiencies — Part 4: Description of causal conditions leading to amputation

	ISO 8548-5:2003
9.	Prosthetics and orthotics — Limb deficiencies — Part 5: Description of the clinical condition of the person who has had an amputation
	ISO 8549-1:2020
10.	Prosthetics and orthotics — Vocabulary — Part 1: General terms for external limb prostheses and external orthoses
	ISO 8549-2:2023
11.	Prosthetics and orthotics — Vocabulary — Part 2: Terms relating to external limb prostheses
	ISO 8549-3:2020
12.	Prosthetics and orthotics — Vocabulary — Part 3: Terms relating to orthoses
	ISO 8549-4:2020
13.	Prosthetics and orthotics — Vocabulary — Part 4: Terms relating to limb amputation
	ISO 8551:2020 5 F D A
14.	Prosthetics and orthotics — Functional deficiencies — Description of the person to be treated with an orthosis, clinical objectives of treatment, and functional requirements of the orthosis
	ISO 13404:2007
15.	Prosthetics and orthotics — Categorization and description of external orthoses and orthotic components
	ISO 13405-1:2015
16.	Prosthetics and orthotics — Classification and description of prosthetic components — Part 1: Classification of prosthetic components
	ISO 13405-2:2015
17.	Prosthetics and orthotics — Classification and description of prosthetic components — Part 2: Description of lower limb prosthetic components

	ISO 13405-3:2015
18.	Prosthetics and orthotics — Classification and description of prosthetic components — Part 3: Description of upper limb prosthetic components
	Description of apper nine prosulette components
	ISO 15032:2000
19.	Prostheses — Structural testing of hip units
	VIO /FIG 1 corr 201 c
20	ISO/TS 16955:2016
20.	Prosthetics — Quantification of physical parameters of ankle foot devices and foot units
	VOO 21062 2017
21	ISO 21063:2017
21.	Prosthetics and orthotics — Soft orthoses — Uses, functions, classification and description
	700 01071 0017
22	ISO 21064:2017
22.	Prosthetics and orthotics — Foot orthotics — Uses, functions classification and description
	ISO 21065-2017
22	ISO 21065:2017
23.	Prosthetics and orthotics — Terms relating to the treatment and rehabilitation of persons having a lower limb amputation
	ISO/TR 22676:2006
24.	Prosthetics — Testing of ankle-foot devices and foot units — Guidance on the application of the test loading conditions of ISO 22675 and on the design of appropriate test equipment
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	ISO 24562:2022
25.	Prosthetics — Geometrical aspects of lower limb prosthetic adapters
	ISO 29781:2008
26.	Prostheses and orthoses — Factors to be included when describing physical activity of a person who has had a lower limb amputation(s) or who has a deficiency of a lower limb segment(s) present at birth
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27.	ISO 29782:2022 Prostheses and orthoses — Factors to be considered when specifying a prosthesis for a person who has had a lower limb amputation
28.	ISO 29783-1:2008 Prosthetics and orthotics — Vocabulary — Part 1: Normal gait
29.	ISO 29783-2:2015 Prosthetics and orthotics — Vocabulary — Part 2: Prosthetic gait
30.	ISO 29783-3:2016 Prosthetics and orthotics — Vocabulary — Part 3: Pathological gait (excluding prosthetic gait)



	Surgical Instruments	
1.	ISO 7151:1988 Surgical instruments — Non-cutting, articulated instruments — General requirements and test methods	
2.	ISO 7153-1:2016 Surgical instruments — Materials — Part 1: Metals	
3.	ISO 7740:1985 Instruments for surgery — Scalpels with detachable blades — Fitting dimensions	
4.	ISO 7741:1986 Instruments for surgery — Scissors and shears — General requirements and test methods	
5.	ISO 13402:1995 Surgical and dental hand instruments — Determination of resistance against autoclaving, corrosion and thermal exposure	

	Sterilization and Disinfectants	
1.	ISO 11135:2014/AMD 1:2018 Sterilization of health-care products — Ethylene oxide — Requirements for the development, validation and routine control of a sterilization process for medical devices — Amendment 1: Revision of Annex E, Single batch release	
2.	ISO 11140-1:2014 Sterilization of health care products - Chemical indicators - Part 1: General requirements	
3.	ISO 11140-3:2007, including Cor 1:2007 Sterilization of health care products - Chemical indicators - Part 3: Class 2 indicator systems for use in the Bowie and Dick-type steam penetration test	
4.	ISO 13408-2:2018 Aseptic processing of health care products - Part 2: Filtration	
5.	ISO 13408-3:2006 Aseptic processing of health care products - Part 3: Lyophilization	
6.	ISO 13408-4:2005 Aseptic processing of health care products - Part 4: Clean-in-place technologies	
7.	ISO 13408-5:2006 Aseptic processing of health care products - Part 5: Sterilization in place	
8.	ISO 13408-7:2012 Aseptic processing of health care products - Part 7: Alternative processes for medical devices and combination products	

9.	ISO 14937:2009 Sterilization of health care products - General requirements for characterization of a sterilizing agent and the development, validation and routine control of a sterilization process for medical devices
10.	ISO 15882:2008 Sterilization of health care products — Chemical indicators — Guidance for selection, use and interpretation of results
11.	ISO 15883-1:2006/Amd 1:2014 Washer-disinfectors - Part 1: General requirements, terms and definitions and tests
12.	ISO 15883-2:2006 Washer-disinfectors - Part 2: Requirements and tests for washer-disinfectors employing thermal disinfection for surgical instruments, anaesthetic equipment, bowls, dishes, receivers, utensils, glassware, etc.
13.	ISO 15883-3:2006 Washer-disinfectors - Part 3: Requirements and tests for washer-disinfectors employing thermal disinfection for human waste containers (ISO 15883-3:2006)
14.	ISO 15883-4:2018 Washer-disinfectors - Part 4: Requirements and tests for washer-disinfectors employing chemical disinfection for thermolabile endoscopes
15.	ISO 15883-5:2021 Washer-disinfectors — Part 5: Performance requirements and test method criteria for demonstrating cleaning efficacy
16.	ISO 15883-6:2011 Washer-disinfectors — Part 6: Requirements and tests for washer-disinfectors employing thermal disinfection for non-invasive, non-critical medical devices and healthcare equipment
17.	ISO 15883-7:2016 Washer-disinfectors — Part 7: Requirements and tests for washer-disinfectors employing chemical disinfection for non-invasive, non-critical thermolabile medical devices and healthcare equipment

18.	ISO/TS 16775:2021 Packaging for terminally sterilized medical devices — Guidance on the application of ISO 11607-1 and ISO 11607-2
19.	ISO 17664-1:2021 Processing of health care products — Information to be provided by the medical device manufacturer for the processing of medical devices — Part 1: Critical and semi-critical medical devices
20.	ISO 17664-2:2021 Processing of health care products — Information to be provided by the medical device manufacturer for the processing of medical devices — Part 2: Non-critical medical devices
21.	ISO 11137-2:2013+Amd 1:2022 Sterilization of health care products - Radiation - Part 2: Establishing the sterilization dose
22.	ISO 11137-3:2017 Sterilization of health care products — Radiation — Part 3: Guidance on dosimetric aspects of development, validation and routine control
23.	ISO/TS 11137-4:2020 Sterilization of health care products — Radiation — Part 4: Guidance on process control
24.	ISO 11138-8:2021 Sterilization of health care products — Biological indicators — Part 8: Method for validation of a reduced incubation time for a biological indicator
25.	ISO 11139:2018 Sterilization of health care products — Vocabulary of terms used in sterilization and related equipment and process standards
26.	ISO 11140-4:2007 Sterilization of health care products — Chemical indicators — Part 4: Class 2 indicators as an alternative to the Bowie and Dick-type test for detection of steam

27.	ISO 11140-5:2007 Sterilization of health care products — Chemical indicators — Part 5: Class 2 indicators for Bowie and Dick-type air removal tests
28.	ISO 11140-6:2022 Sterilization of health care products — Chemical indicators — Part 6: Type 2 indicators and process challenge devices for use in performance testing of small steam sterilizers
29.	ISO 11607-1:2019+Amd 1:2023 Packaging for terminally sterilized medical devices - Part 1: Requirements for materials, sterile barrier systems and packaging systems
30.	ISO 11607-2:2019, including Amd 1:2023 Packaging for terminally sterilized medical devices — Part 2: Validation requirements for forming, sealing and assembly processes
31.	ISO 11737-1:2018 Sterilization of medical devices - Microbiological methods - Part 1: Determination of a population of microorganisms on products
32.	ISO 11737-2:2019 Sterilization of health care products — Microbiological methods — Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process
33.	ISO 11737-3:2023 Sterilization of health care products — Microbiological methods — Part 3: Bacterial endotoxin testing
34.	ISO 13004:2022 Sterilization of health care products — Radiation — Substantiation of selected sterilization dose: Method VDmaxSD
35.	ISO 13408-6:2021 Aseptic processing of health care products — Part 6: Isolator systems

36.	ISO 14160:2020 Sterilization of health care products — Liquid chemical sterilizing agents for single-use medical devices utilizing animal tissues and their derivatives — Requirements for characterization, development, validation and routine control of a sterilization process for medical devices
37.	ISO 18362:2016+Amd 1:2022 Manufacture of cell-based health care products — Control of microbial risks during processing
38.	ISO 18472:2018 Sterilization of health care products — Biological and chemical indicators — Test equipment
39.	ISO/TS 21387:2020 Sterilization of medical devices — Guidance on the requirements for the validation and routine processing of ethylene oxide sterilization processes using parametric release
40.	ISO/TS 22421:2021 Sterilization of health care products — Common requirements for sterilizers for terminal sterilization of medical devices in health care facilities
41.	ISO 22441:2022 Sterilization of health care products — Low temperature vaporized hydrogen peroxide — Requirements for the development, validation and routine control of a sterilization process for medical devices
42.	ISO/TS 22456:2021 Sterilization of healthcare products — Microbiological methods— Guidance on conducting bioburden determinations and tests of sterility for biologics and tissue-based products
43.	ISO 17665-1:2006 Sterilization of health care products moist heat part 1: requirements for the development, validation and routine control of a sterilization process for medical devices
44.	ISO/TS 17665-2:2009 Sterilization of health care products moist heat part 2: guidance on the application of iso 17665-1

45.	ISO/TS 17665-3:2013 Sterilization of health care products moist heat part 3: guidance on the designation of a medical device to a product family and processing category for steam sterilization
46.	ISO 25424:2018+Amd 1:2022 Sterilization of health care products low temperature steam and formaldehyde requirements for development, validation and routine control of a sterilization process for medical devices
47.	EN 13624:2021 Chemical disinfectants and antiseptics — Quantitative suspension test for the evaluation of fungicidal or yeasticidal activity in the medical area — Test method and requirements (phase 2, step 1)
48.	EN 14561:2005 Chemical disinfectants and antiseptics — Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants — Test methods and requirements (phase 2, step 1)
49.	EN 14348:2006 Chemical disinfectants and antiseptics — Quantitative carrier test for the evaluation of bactericidal activity for instruments used in the medical area — Test method and requirements (phase 2, step 2)
50.	ISO/TS 5111:2022 Guidance on quality of water for sterilizers, sterilization and washer-disinfectors for health care products

	In vitro diagnostic	
1.	ISO 18113-1:2022 In vitro diagnostic medical devices Information supplied by the manufacturer (labelling) Part 1: Terms, definitions, and general requirements	
2.	ISO 18113-2:2022 In vitro diagnostic medical devices — Information supplied by the manufacturer (labelling) — Part 2: In vitro diagnostic reagents for professional use	
3.	ISO 18113-3:2022 In vitro diagnostic medical devices — Information supplied by the manufacturer (labelling) — Part 3: In vitro diagnostic instruments for professional use	
4.	ISO 18113-4:2022 In vitro diagnostic medical devices — Information supplied by the manufacturer (labelling) — Part 4: In vitro diagnostic reagents for self-testing	
5.	ISO 18113-5:2022 In vitro diagnostic medical devices — Information supplied by the manufacturer (labelling) — Part 5: In vitro diagnostic instruments for self-testing	
6.	ISO 11137-1:2006 Sterilization of health care products Radiation Part 1: Requirements for development, validation and routine control of a sterilization process for medical devices	
7.	ISO 11137-2:2013 Sterilization of health care products Radiation Part 2: Establishing the sterilization dose	
8.	ISO 11737-1:2018 Sterilization of health care products Microbiological methods Part 1: Determination of a population of microorganisms on products	

9.	ISO 11737-2:2019 Sterilization of health care products Microbiological methods Part 2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process
10.	ISO 11737-3:2023 Sterilization of health care products Microbiological methods Part 3: Bacterial endotoxin testing
11.	ISO 11135:2014 Sterilization of health-care products Ethylene oxide Requirements for the development, validation and routine control of a sterilization process for medical devices
12.	ISO 25424:2018 Sterilization of health care products Low temperature steam and formaldehyde Requirements for development, validation and routine control of a sterilization process for medical devices
13.	ISO 17511:2020 In vitro diagnostic medical devices Requirements for establishing metrological traceability of values assigned to calibrators, trueness control materials and human samples
14.	ISO 6717:2021 In vitro diagnostic medical devices Single-use containers for the collection of specimens from humans other than blood
15.	ISO 15197:2013 In vitro diagnostic test systems Requirements for blood-glucose monitoring systems for self-testing in managing diabetes mellitus
16.	ISO 17511:2020 In vitro diagnostic medical devices Requirements for establishing metrological traceability of values assigned to calibrators, trueness control materials and human samples
17.	ISO 23640:2011 In vitro diagnostic medical devices Evaluation of stability of in vitro diagnostic reagents
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18.	ISO 17593:2022 Clinical laboratory testing and in vitro medical devices Requirements for in vitro monitoring systems for self-testing of oral anticoagulant therapy
19.	ISO 20916:2019 In vitro diagnostic medical devices Clinical performance studies using specimens from human subjects Good study practice
20.	IEC 61010-2-101:2018 Safety requirements for electrical equipment for measurement, control, and laboratory use - Part 2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment
21.	IEC 61326-2-6:2020 Electrical equipment for measurement, control and laboratory use - EMC requirements - Part 2-6: Particular requirements - In vitro diagnostic (IVD) medical equipment
22.	IEC 62366-1:2015/Amd 1:2020 Medical devices Part 1: Application of usability engineering to medical devices
23.	IEC 62304:2006 Medical device software — Software life cycle processes
24.	ISO 13485 :2016 Medical devices Quality management systems Requirements for regulatory purposes
25.	ISO 14971:2019 Medical devices — Application of risk management to medical devices
26.	ISO 20417:2021 Medical devices — Information to be supplied by the manufacturer

27.	ISO 15223-1:2021 Medical devices — Symbols to be used with information to be supplied by the manufacturer — Part 1: General requirements
28.	ISO 15223-2:2010 Medical devices — Symbols to be used with medical device labels, labelling, and information to be supplied — Part 2: Symbol development, selection and validation
29.	ISO/TR 20416:2020 Medical devices — Post-market surveillance for manufacturers
30.	EN 13532:2002 General requirements for in vitro diagnostic medical devices for self-testing
31.	EN 13612:2002 Performance evaluation of in vitro diagnostic medical devices
32.	EN 13641:2002 Elimination or reduction of risk of infection related to in vitro diagnostic reagents

	Electromedical	
1.	IEC 60118-13:2019 Electroacoustics - Hearing aids - Part 13: Requirements and methods of measurement for electromagnetic immunity to mobile digital wireless devices	
2.	IEC 60522-1:2020 Medical electrical equipment - Diagnostics X-rays - Part 1: Determination of quality equivalent filtration and permanent filtration	
3.	IEC TR 60522-2:2020 Medical electrical equipment - Diagnostics X-rays - Part 2: Guidance and rationale on quality equivalent filtration and permanent filtration	
4.	IEC 60580:2019 Medical electrical equipment - Dose area product meters	
5.	IEC 60601-1-2:2014 Medical electrical equipment - Part 1-2: General requirements for basic safety and essential performance - Collateral Standard: Electromagnetic disturbances - Requirements and tests	
6.	IEC 60601-2-1:2009 Medical electrical equipment - Part 2-1: Particular requirements for the safety of electron accelerators in the range of 1 MeV to 50 MeV	
7.	IEC 60601-2-2:2017 Medical electrical equipment - Part 2-2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories	
8.	IEC 60601-2-3:2012+AMD1:2016 CSV Consolidated version Medical electrical equipment - Part 2-3: Particular requirements for the basic safety and essential performance of short-wave therapy equipment	

IEC 60601-2-5:2009
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IEC 60601-2-8:2010+AMD1:2015 CSV Consolidated version
Medical electrical equipment - Part 2-8: Particular requirements for the basic safety and essential performance of therapeutic X-ray equipment operating in the range 10 kV to 1 MV
IEC 60601-2-17:2013
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IEC 60601-2-18:2015
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IEC 60601-2-19:2009+AMD1:2016
Medical electrical equipment - Part 2-19: Particular requirements for the basic safety and essential performance of infant incubators
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IEC 60601-2-27:2011
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18.	IEC 60601-2-28:2017 Medical electrical equipment - Part 2-28: Particular requirements for the basic safety and essential performance of X-ray tube assemblies for medical diagnosis
19.	IEC 60601-2-29:2008 Medical electrical equipment - Part 2-29: Particular requirements for the basic safety and essential performance of radiotherapy simulators
20.	IEC 60601-2-33:2010+AMD1: 2013+AMD2:2015 CSV Consolidated version Medical electrical equipment - Part 2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis
21.	IEC 60601-2-36:2014 Medical electrical equipment - Part 2-36: Particular requirements for the basic safety and essential performance of equipment for extracorporeally induced lithotripsy
22.	IEC 60601-2-37:2007+AMD1:2015 Medical electrical equipment - Part 2-37: Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment
23.	IEC 60601-2-39:2018 Medical electrical equipment - Part 2-39: Particular requirements for basic safety and essential performance of peritoneal dialysis equipment
24.	IEC 60601-2-40:2016 Medical electrical equipment - Part 2-40: Particular requirements for the safety of electromyographs and evoked response equipment
25.	IEC 60601-2-41:2009+AMD1:2013 CSV Consolidated version Medical electrical equipment - Part 2-41: Particular requirements for the basic safety and essential performance of surgical luminaires and luminaires for diagnosis
26.	IEC 60601-2-45:2011+AMD1:2015 CSV Consolidated version Medical electrical equipment - Part 2-45: Particular requirements for the basic safety and essential performance of mammographic X-ray equipment and mammographic stereotactic devices

27.	IEC 60601-2-46:2016 Medical electrical equipment Part 2-46: Particular requirements for the safety of operating tables
28.	IEC 60601-2-47:2012 Medical electrical equipment - Part 2-47: Particular requirements for the safety, including essential performance, of ambulatory electrocardiographic systems
29.	IEC 60601-2-49:2018 Medical electrical equipment - Part 2-49: Particular requirements for the safety of multifunction patient monitoring equipment
30.	IEC 60601-2-50:2009+AMD1:2016 Medical electrical equipment - Part 2-50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment
31.	IEC 60601-2-52:2009 Medical electrical equipment Part 2-52: Particular requirements for the basic safety and essential performance of medical beds
32.	IEC 60601-2-52:2009/Cor 1:2010 Medical electrical equipment Part 2-52: Particular requirements for the basic safety and essential performance of medical beds Technical Corrigendum 1
33.	IEC 60601-2-52:2009/Amd 1:2015 Medical electrical equipment Part 2-52: Particular requirements for the basic safety and essential performance of medical beds Amendment 1
34.	IEC 60645-1:2017 Electroacoustic - Audiological equipment - Part 1: Pure-tone audiometers

25	IEC 61676:2002+AMD1:2008
35.	Medical electrical equipment - Dosimetric instruments used for non-invasive measurement of X-ray tube voltage in diagnostic radiology
	IEC 62220-1-1:2015
36.	Medical electrical equipment - Characteristics of digital X-ray imaging devices - Part 1-1: Determination of the detective quantum efficiency - Detectors used in radiographic imaging
	IEC 62220-1-2:2007
37.	Medical electrical equipment - Characteristics of digital X-ray imaging devices - Part 1-2: Determination of the detective quantum efficiency - Detectors used in mammography
	IEC 62220-1-3:2008
38.	Medical electrical equipment - Characteristics of digital X-ray imaging devices - Part 1-3: Determination of the detective quantum efficiency - Detectors used in dynamic imaging
	IEC 80601-2-35:2009+AMD1:2016
39.	Medical electrical equipment - Part 2-35: Particular requirements for the basic safety and essential performance of heating devices using blankets, pads and mattresses and intended for heating in medical use
	IEC 80601-2-31:2020
40.	Medical electrical equipment - Part 2-31: Particular requirements for the basic safety and essential performance of external cardiac pacemakers with internal power source
	IEC 80601-2-58:2014+AMD1:2016 CSV Consolidated version
41.	Medical electrical equipment - Part 2-58: Particular requirements for the basic safety and essential performance of lens removal devices and vitrectomy devices for ophthalmic surgery
	IEC 80601-2-59:2017
42.	Medical electrical equipment - Part 2-59: Particular requirements for the basic safety and essential performance of screening thermographs for human febrile temperature screening
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44.	IEC 60601-1:2005+AMD1:2012 Medical electrical equipment - Part 1: General requirements for basic safety and essential performance
45.	IEC 60601-1-6:2010+AMD1:2013+AMD2:2020 Medical electrical equipment - Part 1-6: General requirements for basic safety and essential performance - Collateral standard: Usability
46.	IEC 80601-2-26:2019 Medical electrical equipment - Part 2-26: Particular requirements for the basic safety and essential performance of electroencephalograph
47.	IEC 60601-2-31:2020 Medical electrical equipment - Part 2-31: Particular requirements for the basic safety and essential performance of external cardiac pacemakers with internal power source
48.	IEC 60601-2-57:2011 Medical electrical equipment – Part 2-57: Particular requirements for the basic safety and essential performance of non-laser light source equipment intended for therapeutic, diagnostic, monitoring and cosmetic/aesthetic use
49.	ISO 81060-2:2018 Non-invasive sphygmomanometers — Part 2: Clinical investigation of intermittent automated measurement type
50.	ISO 81060-2:2018/Amd 1:2020 Non-invasive sphygmomanometers — Part 2: Clinical investigation of intermittent automated measurement type — Amendment 1
51.	IEC 60601-2-4/AMD 2 Amendment 2 - Medical electrical equipment - Part 2-4: Particular requirements for the basic safety and essential performance of cardiac defibrillators

52.	ISO 81060-3:2022 Non-invasive sphygmomanometers — Part 3: Clinical investigation of continuous automated measurement type
53.	ISO/TS 81060-5:2020 Non-invasive sphygmomanometers — Part 5: Requirements for the repeatability and reproducibility of NIBP simulators for testing of automated non-invasive sphygmomanometers
54.	ISO 80601-2-56:2017 Medical electrical equipment — Part 2-56: Particular requirements for basic safety and essential performance of clinical thermometers for body temperature measurement
55.	ISO 80601-2-56:2017/Amd 1:2018 Medical electrical equipment — Part 2-56: Particular requirements for basic safety and essential performance of clinical thermometers for body temperature measurement — Amendment 1
56.	ISO 80601-2-80:2018 Medical electrical equipment - Part 2-80: Particular requirements for basic safety and essential performance of ventilatory support equipment for ventilatory insufficiency
57.	ISO 80601-2-90:2021 Medical electrical equipment Part 2-90: Particular requirements for basic safety and essential performance of respiratory high-flow therapy equipment
58.	IEC-62281:2016 Safety of primary and secondary lithium cells and batteries during transport
59.	ISO 80601-2-90:2021 Medical electrical equipment — Part 2-90: Particular requirements for basic safety and essential performance of respiratory high-flow therapy equipment
60.	IEC-60601-1-10:2015

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61.	IEC 80601-2-49:2018 Medical electrical equipment — Part 2-49: Particular requirements for the basic safety and essential performance of multifunction patient monitoring equipment
62.	IEC 80601-2-52 Medical electrical equipment - Part 2-52: Particular requirements for the basic safety and essential performance of medical beds
63.	IEC 80601-2-71:2015 Medical electrical equipment — Part 2-71: Particular requirements for the basic safety and essential performance of functional Near-Infrared Spectroscopy (NIRS) equipment
64.	ISO 80601-2-85:2021 Medical electrical equipment — Part 2-85: Particular requirements for the basic safety and essential performance of cerebral tissue oximeter equipment
65.	IEC 80601-2-30:2018 Medical electrical equipment — Part 2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers
66.	ISO 80601-2-61:2017 Medical electrical equipment — Part 2-61: Particular requirements for basic safety and essential performance of pulse oximeter equipment

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2.	IEC 62220-1:2015 Medical electrical equipment - Characteristics of digital X-ray imaging devices - Part 1-1: Determination of the detective quantum efficiency - Detectors used in radiographic imaging
3.	IEC 60627:2013 Diagnostic X-ray imaging equipment - Characteristics of general purpose and mammographic antiscatter grids
4.	ISO 15382:2015 Radiological protection — Procedures for monitoring the dose to the lens of the eye, the skin and the extremities
5.	ISO 5576:1997 Non-destructive testing — Industrial X-ray and gamma-ray radiology — Vocabulary
6.	ISO 5579:2013 Non-destructive testing — Radiographic testing of metallic materials using film and X- or gamma rays — Basic rules
7.	ISO 5580:1985 Non-destructive testing — Industrial radiographic illuminators — Minimum requirements
8.	ISO 11699-1:2008 Non-destructive testing — Industrial radiographic film — Part 1: Classification of film systems for industrial radiography
9.	ISO 11699-2:2018 Non-destructive testing — Industrial radiographic films — Part 2: Control of film processing by means of reference values

10.	ISO 12721:2000 Non-destructive testing — Thermal neutron radiographic testing — Determination of beam L/D ratio
11.	ISO 14096-1:2005 Non-destructive testing — Qualification of radiographic film digitisation systems — Part 1: Definitions, quantitative measurements of image quality parameters, standard reference film and qualitative control
12.	ISO 14096-2:2005 Non-destructive testing — Qualification of radiographic film digitisation systems — Part 2: Minimum requirements
13.	ISO 15708-1:2017 Non-destructive testing — Radiation methods for computed tomography — Part 1: Terminology
14.	ISO 15708-2:2017 Non-destructive testing — Radiation methods for computed tomography — Part 2: Principles, equipment and samples
15.	ISO 15708-3:2017 Non-destructive testing — Radiation methods for computed tomography — Part 3: Operation and interpretation
16.	ISO 15708-4:2017 Non-destructive testing — Radiation methods for computed tomography — Part 4: Qualification
17.	ISO 16371-1:2011 Non-destructive testing — Industrial computed radiography with storage phosphor imaging plates — Part 1: Classification of systems
18.	ISO 16371-2:2017 Non-destructive testing — Industrial computed radiography with storage phosphor imaging plates — Part 2: General principles for testing of metallic materials using X-rays and gamma rays

19.	ISO 16526-1:2011 Non-destructive testing — Measurement and evaluation of the X-ray tube voltage — Part 1: Voltage divider method
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21.	ISO 16526-3:2011 Non-destructive testing — Measurement and evaluation of the X-ray tube voltage — Part 3: Spectrometric method
22.	ISO 19232-1:2013 Non-destructive testing — Image quality of radiographs — Part 1: Determination of the image quality value using wire-type image quality indicators
23.	ISO 19232-2:2013 Non-destructive testing — Image quality of radiographs — Part 2: Determination of the image quality value using step/hole-type image quality indicators
24.	ISO 19232-3:2013 Non-destructive testing — Image quality of radiographs — Part 3: Image quality classes
25.	ISO 19232-4:2013 Non-destructive testing — Image quality of radiographs — Part 4: Experimental evaluation of image quality values and image quality tables
26.	ISO 19232-5:2018 Non-destructive testing — Image quality of radiographs — Part 5: Determination of the image unsharpness and basic spatial resolution value using duplex wire-type image quality indicators
27.	ISO 20769-1:2018 Non-destructive testing — Radiographic inspection of corrosion and deposits in pipes by X- and gamma rays — Part 1: Tangential radiographic inspection

28.	ISO 20769-2:2018 Non-destructive testing — Radiographic inspection of corrosion and deposits in pipes by X- and gamma rays — Part 2: Double wall radiographic inspection
29.	ISO 21432:2019 Non-destructive testing — Standard test method for determining residual stresses by neutron diffraction
30.	ISO 23159:2020 Non-destructive testing — Gamma ray scanning method on process columns



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1.	ISO 3990:2023 Dentistry — Evaluation of antibacterial activity of dental restorative materials, luting materials, fissure sealants and orthodontic bonding or luting materials	
2.	ISO 7405:2018 Dentistry — Evaluation of biocompatibility of medical devices used in dentistry	
3.	ISO 3107:2022 Dentistry — Zinc oxide-eugenol cements and non-eugenol zinc oxide cements	
4.	ISO 4049:2019 Dentistry — Polymer-based restorative materials	
5.	ISO/TS 4640:2023 Dentistry — Test methods for tensile bond strength to tooth structure	
6.	ISO 6872:2015+ Amd 1:2018 Dentistry — Ceramic materials	
7.	ISO 6874:2015 Dentistry — Polymer-based pit and fissure sealants	
8.	ISO 6876:2012 Dentistry — Root canal sealing materials	
9.	ISO 6877:2021 Dentistry — Endodontic obturating materials	
10.	ISO 7405:2018 Dentistry — Evaluation of biocompatibility of medical devices used in dentistry	

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11.	
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12.	ISO 9917-1:2007
	Dentistry — Water-based cements — Part 1: Powder/liquid acid-base cements
	Dentistry Water-based cements Tart 1. Towaer/inquita acid-base cements
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13.	Dentistry — Water-based cements — Part 2: Resin-modified cements
14	ISO 9693:2019
14.	Dentistry — Compatibility testing for metal-ceramic and ceramic-ceramic systems
	ISO 10271:2020
15.	
	Dentistry — Corrosion test methods for metallic materials
	ISO 14801:2016
16.	Dentistry — Implants — Dynamic loading test for endosseous dental implants
	SEDA
17	ISO 22674:2022
17.	Dentistry — Metallic materials for fixed and removable restorations and appliances
	ISO 22794:2007
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18.	Dentistry — Implantable materials for bone filling and augmentation in oral and maxillofacial surgery — Contents of a technical file
	ISO 22803:2004
19.	Dentistry — Membrane materials for guided tissue regeneration in oral and maxillofacial surgery —
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20.	Dentistry — Dental amalgam

21.	ISO/TS 11405:2015 Dentistry — Testing of adhesion to tooth structure
22.	ISO 13116:2014 Dentistry — Test method for determining radio-opacity of materials
23.	ISO 29022:2013 Dentistry — Adhesion — Notched-edge shear bond strength test
24.	ISO 6873:2013 Dentistry — Gypsum products
25.	ISO 7491:2000 Dental materials — Determination of colour stability
26.	ISO 9333:2022 Dentistry — Brazing materials
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28.	ISO 10139-1:2018 Dentistry — Soft lining materials for removable dentures — Part 1: Materials for short-term use
29.	ISO 10139-2:2016 Dentistry — Soft lining materials for removable dentures — Part 2: Materials for long-term use
30.	ISO 10271:2020 Dentistry — Corrosion test methods for metallic materials
31.	ISO 10477:2020 Dentistry — Polymer-based crown and veneering materials

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32.	Dentistry — Magnetic attachments
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33.	Dentistry — Dental furnace — Part 2: Test method for evaluation of furnace programme via firing glaze
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34.	Dentistry — Dental furnace — Part 3: Test method for the evaluation of high temperature sintering furnace measurement with a separate thermocouple
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27	ISO 14356:2003
37.	Dentistry — Duplicating material 5 F D A
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38.	Dental materials — Guidance on testing of wear — Part 1: Wear by toothbrushing
20	ISO/TS 14569-2:2001
39.	Dental materials — Guidance on testing of wear — Part 2: Wear by two- and/or three body contact
40.	ISO 15854:2023
	Dentistry — Casting and baseplate waxes
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41.	Dentistry — Bonding test between polymer teeth and denture base materials

42.	ISO 20795-1:2013 Dentistry — Base polymers — Part 1: Denture base polymers
43.	ISO 20795-2:2013 Dentistry — Base polymers — Part 2: Orthodontic base polymers
44.	ISO 21563:2021 Dentistry — Hydrocolloid impression materials
45.	ISO 22112:2017 Dentistry — Artificial teeth for dental prostheses
46.	ISO 22598:2020 Dentistry — Colour tabs for intraoral tooth colour determination
47.	ISO 22674:2022 Dentistry — Metallic materials for fixed and removable restorations and appliances
48.	ISO 23401-1:2023 Dentistry — Chairside denture base relining materials — Part 1: Hard type materials
49.	ISO 28319:2018 Dentistry — Laser welding and filler materials
50.	ISO 1797:2017 Dentistry — Shanks for rotary and oscillating instruments
51.	ISO 3630-1:2019 Dentistry — Endodontic instruments — Part 1: General requirements
52.	ISO 3630-2:2023 Dentistry — Endodontic instruments — Part 2: Enlargers

53.	ISO 3630-3:2021 Dentistry — Endodontic instruments — Part 3: Compactors
54.	ISO 3630-4:2023 Dentistry — Endodontic instruments — Part 4: Auxiliary instruments
55.	ISO 3630-5:2020 Dentistry — Endodontic instruments — Part 5: Shaping and cleaning instruments
56.	ISO/TR 3630-6:2023 Dentistry — Endodontic instruments — Part 6: Numeric coding system
57.	ISO 3823-1:1997 Dental rotary instruments — Burs — Part 1: Steel and carbide burs
58.	ISO 3823-2:2003+ Amd 1:2008 Dentistry — Rotary bur instruments — Part 2: Finishing burs
59.	ISO 3964:2016+ Amd 1:2018 Dentistry — Coupling dimensions for handpiece connectors
60.	ISO 4865-1:2023 Dentistry — General requirements of hand instruments — Part 1: Non-hinged hand instruments
61.	ISO 6360-1:2004+ Cor 1:2007 Dentistry — Number coding system for rotary instruments — Part 1: General characteristics
62.	ISO 6360-2:2004+ Amd 1:2011 Dentistry — Number coding system for rotary instruments — Part 2: Shapes
63.	ISO 6360-3:2005

	Dentistry — Number coding system for rotary instruments — Part 3: Specific characteristics of burs and cutters
64.	ISO 6360-4:2004 Dentistry — Number coding system for rotary instruments — Part 4: Specific characteristics of diamond instruments
65.	ISO 6360-5:2007 Dentistry — Number coding system for rotary instruments — Part 5: Specific characteristics of root-canal instruments
66.	ISO 6360-6:2004 Dentistry — Number coding system for rotary instruments — Part 6: Specific characteristics of abrasive instruments
67.	ISO 6360-7:2006 Dentistry — Number coding system for rotary instruments — Part 7: Specific characteristics of mandrels and special instruments
68.	ISO 7492:2019 Dentistry — Dental explorer
69.	ISO 7711-1:2021 Dentistry — Diamond rotary instruments — Part 1: General requirements
70.	ISO 7711-2:2011 Dentistry — Rotary diamond instruments — Part 2: Discs
71.	ISO 7786:2001 Dental rotary instruments — Laboratory abrasive instruments
72.	ISO 7787-1:2016 Dentistry — Laboratory cutters — Part 1: Steel laboratory cutters

73.	ISO 7787-2:2020 Dentistry — Laboratory cutters — Part 2: Carbide laboratory cutters
74.	ISO 7787-3:2017 Dentistry — Laboratory cutters — Part 3: Carbide cutters for milling machines
75.	ISO 7787-4:2002 Dental rotary instruments — Cutters — Part 4: Miniature carbide laboratory cutters
76.	ISO 7885:2010 Dentistry — Sterile injection needles for single use
77.	ISO 8325:2023 Dentistry — Test methods for rotary instruments
78.	ISO 9168:2009 Dentistry — Hose connectors for air driven dental handpieces
79.	ISO 9173-1:2016 Dentistry — Extraction forceps — Part 1: General requirements
80.	ISO 9173-2:2010 Dentistry — Extraction forceps — Part 2: Designation
81.	ISO 9173-3:2014 Dentistry — Extraction forceps — Part 3: Design
82.	ISO 9873:2019 Dentistry — Intra-oral mirrors
83.	ISO 9997:2020 Dentistry — Cartridge syringes

84.	ISO 10323:2013
	Dentistry — Bore diameters for rotary instruments such as discs and wheels
85.	ISO 11499:2014
	Dentistry — Single-use cartridges for local anaesthetics
86.	ISO 13295:2007
	Dentistry — Mandrels for rotary instruments
87.	ISO 13397-1:1995
	Periodontal curettes, dental scalers and excavators — Part 1: General requirements
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88.	Dentistry — Periodontal curettes, dental scalers and excavators — Part 2: Periodontal curettes of Grtype
89.	ISO 13397-3:1996
09.	Periodontal curettes, dental scalers and excavators — Part 3: Dental scalers — H-type
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90.	Dentistry — Periodontal curettes, dental scalers and excavators — Part 5: Jacquette scalers
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91.	Dentistry — General requirements for instruments and related accessories used in dental implant placement and treatment
	ISO 14457:2017
92.	Dentistry — Handpieces and motors
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93.	Dental elevators — Part 1: General requirements

94.	ISO 15087-2:2000 Dental elevators — Part 2: Warwick James elevators
95.	ISO 15087-3:2000 Dental elevators — Part 3: Cryer elevators
96.	ISO 15087-4:2000 Dental elevators — Part 4: Coupland elevators
97.	ISO 15087-5:2000 Dental elevators — Part 5: Bein elevators
98.	ISO 15087-6:2000 Dental elevators — Part 6: Flohr elevators
99.	ISO 15098:2020 Dentistry — Dental tweezers
100.	ISO 16635-1:2013 Dentistry — Dental rubber dam technique — Part 1: Hole punch
101.	ISO 16635-2:2014 Dentistry — Dental rubber dam instruments — Part 2: Clamp forceps
102.	ISO 17509:2016 Dentistry — Torque transmitter for handpieces
103.	ISO 17937:2015 Dentistry — Osteotome
104.	ISO 18397:2016 Dentistry — Powered scaler

105.	ISO 18556:2016
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106.	ISO 19490:2017 Dentistry — Sinus membrane elevator
107.	ISO 19715:2017 Dentistry — Filling instrument with contra angle
100	ISO 20569:2018
108.	Dentistry — Trephine burs
109.	ISO 20570:2018
109.	Dentistry — Oral surgical scalpel handle
110.	ISO 20608:2018
110.	Dentistry — Powder jet handpieces and powders
111.	ISO 21531:2009
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112.	ISO 21533:2018
	Dentistry — Reprocessable cartridge syringes for intraligamentary injections
113.	ISO 21671:2006+Amd 1:2011
	Dentistry — Rotary polishers
114.	ISO 21672-1:2012
	Dentistry — Periodontal probes — Part 1: General requirements
115.	ISO 21672-2:2012
	Dentistry — Periodontal probes — Part 2: Designation

116.	ISO 21850-1:2020 Dentistry — Materials for dental instruments — Part 1: Stainless steel
117.	ISO 22569:2020 Dentistry — Multifunction handpieces
118.	ISO 22570:2020 Dentistry — Spoons and bone curettes
119.	ISO 23445:2021 Dentistry — Tissue punches
120.	ISO 23450:2021 Dentistry — Intraoral camera
121.	ISO 23940:2021 Dentistry — Excavators
122.	ISO 5467-1:2022 Dentistry — Mobile dental units and dental patient chairs — Part 1: General requirements
123.	ISO 7488:2018 Dentistry — Mixing machines for dental amalgam
124.	ISO 7493:2006 Dentistry — Operator's stool
125.	ISO 7494-1:2018 Dentistry — Stationary dental units and dental patient chairs — Part 1: General requirements
126.	ISO 7494-2:2022 Dentistry — Stationary dental units and dental patient chairs — Part 2: Air, water, suction and wastewater systems

127.	ISO 9680:2021 Dentistry — Operating lights
128.	ISO 9687:2015+ Amd 1:2018 Dentistry — Graphical symbols for dental equipment
129.	ISO 10637:2018 Dentistry — Central suction source equipment
130.	ISO 10650:2018 Dentistry — Powered polymerization activators
131.	ISO 11143:2008 Dentistry — Amalgam separators
132.	ISO 13897:2018 Dentistry — Dental amalgam reusable mixing-capsules
133.	ISO 16954:2015 Dentistry — Test methods for dental unit waterline biofilm treatment
134.	ISO 21530:2004 Dentistry — Materials used for dental equipment surfaces — Determination of resistance to chemical disinfectants
135.	ISO 22052:2020 Dentistry — Central compressed air source equipment
136.	ISO 23402-1:2020 Dentistry — Portable dental equipment for use in non-permanent healthcare environment — Part 1: General requirements
137.	IEC 80601-2-60:2019

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138.	ISO 10451:2010 Dentistry — Contents of technical file for dental implant systems
139.	ISO 11953:2010 Dentistry — Implants — Clinical performance of hand torque instruments
140.	ISO/TS 13498:2011 Dentistry — Torsion test of implant body/connecting part joints of endosseous dental implant systems
141.	ISO 14801:2016 Dentistry — Implants — Dynamic loading test for endosseous dental implants
142.	ISO 16498:2013 Dentistry — Minimal dental implant data set for clinical use
143.	ISO/TR 18130:2016 Dentistry — Screw loosening test using cyclic torsional loading for implant body/implant abutment connection of endosseous dental implants
144.	ISO 19429:2015 Dentistry — Designation system for dental implants
145.	ISO 22683:2022 Dentistry — Rotational adaptability test between implant body and implant abutment in dental implant system
146.	ISO 22794:2007 Dentistry — Implantable materials for bone filling and augmentation in oral and maxillofacial surgery — Contents of a technical file

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147.	Dentistry — Membrane materials for guided tissue regeneration in oral and maxillofacial surgery — Contents of a technical file
148.	ISO 5139:2023 Dentistry — Polymer-based composite machinable blanks
149.	ISO 12836:2015 Dentistry — Digitizing devices for CAD/CAM systems for indirect dental restorations — Test methods for assessing accuracy
150.	ISO 18618:2022 Dentistry — Interoperability of CAD/CAM systems
151.	ISO 18675:2022 Dentistry — Machinable ceramic blanks
152.	ISO 20896-1:2019 Dentistry — Digital impression devices — Part 1: Methods for assessing accuracy
153.	ISO/TR 20896-2:2023 Dentistry — Digital impression devices — Part 2: Methods for assessing accuracy for implanted devices
154.	ISO/TR 22710:2019 Dentistry — Vocabulary of process chain from dental CT to CAD/CAM for implant prosthetic restorations — Backward planning in the digital process chain
155.	ISO 23298:2023 Dentistry — Test methods for machining accuracy of computer-aided milling machines

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1.	ISO 7998:2005 Ophthalmic optics — Spectacle frames — Lists of equivalent terms and vocabulary	
2.	ISO 8429:1986 Optics and optical instruments — Ophthalmology — Graduated dial scale	
3.	ISO 8596:2017 Ophthalmic optics — Visual acuity testing — Standard and clinical optotypes and their presentation	
4.	ISO 8596:2017/Amd 1:2019 Ophthalmic optics — Visual acuity testing — Standard and clinical optotypes and their presentation — Amendment 1	
5.	ISO 8598-1:2014 Optics and optical instruments — Focimeters — Part 1: General purpose instruments	
6.	ISO 8612:2009 Ophthalmic instruments — Tonometers	
7.	ISO 8624:2020 Ophthalmic optics — Spectacle frames — Measuring system and vocabulary	
8.	ISO 8980-1:2017 Ophthalmic optics — Uncut finished spectacle lenses — Part 1: Specifications for single-vision and multifocal lenses	
9.	ISO 8980-2:2017 Ophthalmic optics — Uncut finished spectacle lenses — Part 2: Specifications for power-variation lenses	

10.	ISO 8980-3:2022 Ophthalmic optics — Uncut finished spectacle lenses — Part 3: Transmittance specifications and test methods
11.	ISO 8980-4:2006 Ophthalmic optics — Uncut finished spectacle lenses — Part 4: Specifications and test methods for anti-reflective coatings
12.	ISO 8980-5:2005 Ophthalmic optics — Uncut finished spectacle lenses — Part 5: Minimum requirements for spectacle lens surfaces claimed to be abrasion-resistant
13.	ISO 9342-1:2023 Optics and optical instruments — Test lenses for calibration of focimeters — Part 1: Reference lenses for focimeters used for measuring spectacle lenses
14.	ISO 9342-2:2005 Optics and optical instruments — Test lenses for calibration of focimeters — Part 2: Test lenses for focimeters used for measuring contact lenses
15.	ISO 9394:2012 Ophthalmic optics — Contact lenses and contact lens care products — Determination of biocompatibility by ocular study with rabbit eyes
16.	ISO 9801:2009 Ophthalmic instruments — Trial case lenses
17.	ISO 10322-1:2016 Ophthalmic optics — Semi-finished spectacle lens blanks — Part 1: Specifications for single-vision and multifocal lens blanks
18.	ISO 10341:2012 Ophthalmic instruments — Refractor heads

19.	ISO 10342:2010 Ophthalmic instruments — Eye refractometers
20.	ISO 10343:2014 Ophthalmic instruments — Ophthalmometers
21.	ISO 10939:2017 Ophthalmic instruments — Slit-lamp microscopes
22.	ISO 10940:2009 Ophthalmic instruments — Fundus cameras
23.	ISO 10942:2022 Ophthalmic instruments — Direct ophthalmoscopes
24.	ISO 10943:2023 Ophthalmic instruments — Indirect ophthalmoscopes
25.	ISO 10944:2009 Ophthalmic instruments — Synoptophores
26.	ISO 11978:2017 Ophthalmic optics — Contact lenses and contact lens care products — Labelling
27.	ISO 11978:2017/Amd 1:2020 Ophthalmic optics — Contact lenses and contact lens care products — Labelling — Amendment 1

28.	ISO 11979-1:2018 Ophthalmic implants — Intraocular lenses — Part 1: Vocabulary
29.	ISO 11979-2:2014 Ophthalmic implants — Intraocular lenses — Part 2: Optical properties and test methods
30.	ISO 11979-3:2012 Ophthalmic implants — Intraocular lenses — Part 3: Mechanical properties and test methods
31.	ISO 11979-4:2008 Ophthalmic implants — Intraocular lenses — Part 4: Labelling and information
32.	ISO 11979-4:2008/Amd 1:2012 Ophthalmic implants — Intraocular lenses — Part 4: Labelling and information — Amendment 1
33.	ISO 11979-5:2020 Ophthalmic implants — Intraocular lenses — Part 5: Biocompatibility
34.	ISO 11979-6:2014 Ophthalmic implants — Intraocular lenses — Part 6: Shelf-life and transport stability testing
35.	ISO 11979-7:2018 Ophthalmic implants — Intraocular lenses — Part 7: Clinical investigations of intraocular lenses for the correction of aphakia
36.	ISO 11979-8:2017 Ophthalmic implants — Intraocular lenses — Part 8: Fundamental requirements

37.	ISO 11981:2017 Ophthalmic optics — Contact lenses and contact lens care products — Determination of physical compatibility of contact lens care products with contact lenses
38.	ISO 11987:2012 Ophthalmic optics — Contact lenses — Determination of shelf-life
39.	ISO 12865:2006 Ophthalmic instruments — Retinoscopes
40.	ISO 12867:2010 Ophthalmic instruments — Trial frames
41.	ISO 14534:2011 Ophthalmic optics — Contact lenses and contact lens care products — Fundamental requirements
42.	ISO 14729:2001 Ophthalmic optics — Contact lens care products — Microbiological requirements and test methods for products and regimens for hygienic management of contact lenses
43.	ISO 14730:2014 Ophthalmic optics — Contact lens care products — Antimicrobial preservative efficacy testing and guidance on determining discard date
44.	ISO 14889:2013/Amd 1:2017 Ophthalmic optics — Spectacle lenses — Fundamental requirements for uncut finished lenses — Amendment 1
45.	ISO 15253:2021 Ophthalmic optics and instruments — Optical and electro-optical devices for enhancing low vision

46.	ISO 15798:2022 Ophthalmic implants — Ophthalmic viscosurgical devices
47.	ISO 16034:2002 Ophthalmic optics — Specifications for single-vision ready-to-wear near- vision spectacles
48.	ISO 16672:2020 Ophthalmic implants — Ocular endotamponades
49.	ISO 16971:2015 Ophthalmic instruments — Optical coherence tomograph for the posterior segment of the human eye
50.	ISO 18189:2016 Ophthalmic optics — Contact lenses and contact lens care products — Cytotoxicity testing of contact lenses in combination with lens care solution to evaluate lens/solution interactions
51.	ISO 18259:2014 Ophthalmic optics — Contact lens care products — Method to assess contact lens care products with contact lenses in a lens case, challenged with bacterial and fungal organisms
52.	ISO 18369-1:2017 Ophthalmic optics — Contact lenses — Part 1: Vocabulary, classification system and recommendations for labelling specifications
53.	ISO 18369-3:2017 Ophthalmic optics — Contact lenses — Part 3: Measurement methods
54.	ISO 18369-4:2017 Ophthalmic optics — Contact lenses — Part 4: Physicochemical properties of contact lens materials

55.	ISO 19045:2015 Ophthalmic optics — Contact lens care products — Method for evaluating Acanthamoeba encystment by contact lens care products
56.	ISO 19980:2021 Ophthalmic instruments — Corneal topographers
57.	ISO 21987:2017 Ophthalmic optics — Mounted spectacle lenses
58.	ISO 22665:2012 Ophthalmic optics and instruments - Instruments to measure axial distances in the eye

SFDA

	Health Informatics
1.	IEC/TR 80001-2-3:2012 Application of risk management for IT-networks incorporating medical devices — Part 2-3: Guidance for wireless networks
2.	IEC/TR 80001-2-4:2012 Application of risk management for IT-networks incorporating medical devices — Part 2-4: General implementation guidance for Healthcare Delivery Organizations
3.	IEC/TR 80001-2-5:2014 Application of risk management for IT-networks incorporating medical devices — Part 2-5: Application guidance — Guidance for distributed alarm systems
4.	IEC/TR 80001-2-8:2016 Application of risk management for IT-networks incorporating medical devices — Part 2-8: Application guidance — Guidance on standards for establishing the security capabilities identified in IEC 80001-2-2
5.	IEC/TR 80001-2-9:2017 Application of risk management for IT-networks incorporating medical devices — Part 2-9: Application guidance — Guidance for use of security assurance cases to demonstrate confidence in IEC/TR 80001-2-2 security capabilities
6.	IEC/TR 80001-2-6:2014 Application of risk management for IT-networks incorporating medical devices — Part 2-6: Application guidance — Guidance for responsibility agreements
7.	IEC/TR 80001-2-7:2015 Application of risk management for IT-networks incorporating medical devices — Application guidance — Part 2-7: Guidance for healthcare delivery organizations (HDOs) on how to self-assess their conformance with IEC 80001-1
8.	IEC 82304-1:2016 Health software — Part 1: General requirements for product safety

	Assistive Products	
1.	ISO 19894:2019 Walking trolleys — Requirements and test methods	
2.	ISO 7176-1:2014 Wheelchairs — Part 1: Determination of static stability	
3.	ISO 7176-5:2008 Wheelchairs — Part 5: Determination of dimensions, mass and manoeuvring space	
4.	ISO 7176-8:2014 Wheelchairs part 8: requirements and test methods for static, impact and fatigue strengths	
5.	ISO 7176-11:2012 Wheelchairs — Part 11: Test dummies 5 F D A	
6.	ISO 7176-13:1989 Wheelchairs — Part 13: Determination of coefficient of friction of test surfaces	
7.	ISO 7176-15:1996 Wheelchairs — Part 15: Requirements for information disclosure, documentation and labelling	
8.	ISO 7176-16:2012 Wheelchairs part 16: resistance to ignition of postural support devices	
9.	ISO 7176-19:2008/AMD 1:2015 Wheelchairs — Part 19: Wheeled mobility devices for use as seats in motor vehicles- Amendment 1: Annex G	

	ISO 7176-21:2009
10.	Wheelchairs part 21: requirements and test methods for electromagnetic compatibility of electrically powered wheelchairs and scooters, and battery chargers
	ISO 7176-25:2022
11.	Wheelchairs Part 25: Lead-acid batteries and chargers for powered wheelchairs. Requirements and test methods
	ISO 21856:2022
12.	Assistive products — General requirements and test methods
	ISO/TR 11548-1:2001
13.	Communication aids for blind persons — Identifiers, names and assignation to coded character sets for 8-dot Braille characters — Part 1: General guidelines for Braille identifiers and shift marks
	ISO/TR 11548-2:2001
14.	Communication aids for blind persons — Identifiers, names and assignation to coded character sets for 8-dot Braille characters — Part 2: Latin alphabet based character sets
	ISO 20342-1:2022
15.	Assistive products for tissue integrity when lying down — Part 1: General requirements
	ISO/TR 20342-7:2021
16.	Assistive products for tissue integrity when lying down — Part 7: Foam properties, characteristics and performance
	ISO 21801-1:2020
17.	Cognitive accessibility — Part 1: General guidelines
	ISO 21801-2:2022
18.	Cognitive accessibility — Part 2: Reporting

	ISO 21802:2019
19.	Assistive products — Guidelines on cognitive accessibility — Daily time management
	ISO 23600:2007
20.	Assistive products for persons with vision impairments and persons with vision and hearing impairments — Acoustic and tactile signals for pedestrian traffic lights
	ISO 24415-2:2011
21.	Tips for assistive products for walking — Requirements and test methods — Part 2: Durability of tips for crutches
	ISO 7176-2:2017
22.	Wheelchairs — Part 2: Determination of dynamic stability of electrically powered wheelchairs
	ISO 7176-4:2008
23.	Wheelchairs — Part 4: Energy consumption of electric wheelchairs and scooters for determination of theoretical distance range
	ISO 7176-6:2018
24.	Wheelchairs — Part 6: Determination of maximum speed of electrically powered wheelchairs
	ISO 7176-10:2008
25.	Wheelchairs — Part 10: Determination of obstacle-climbing ability of electrically powered wheelchairs
	ISO 7176-22:2014
26.	Wheelchairs — Part 22: Set-up procedures
	ISO 7176-26:2007
27.	Wheelchairs — Part 26: Vocabulary

	ISO 7176-28:2012
28.	Wheelchairs — Part 28: Requirements and test methods for stair-climbing devices
	ISO 7176-30:2018
29.	Wheelchairs — Part 30: Wheelchairs for changing occupant posture — Test methods and requirements
	ISO 7176-31:2023
30.	Wheelchairs — Part 31: Lithium-ion battery systems and chargers for powered wheelchairs — Requirements and test methods
	ISO 7176-32:2022
31.	Wheelchairs — Part 32: Test method for wheelchair castor assembly durability
	ISO 10542-1:2012
32.	Technical systems and aids for disabled or handicapped persons — Wheelchair tiedown and occupant-restraint systems — Part 1: Requirements and test methods for all systems
	ISO 10542-1:2012/Amd 1:2021
33.	Technical systems and aids for disabled or handicapped persons — Wheelchair tiedown and occupant-restraint systems — Part 1: Requirements and test methods for all systems — Amendment 1: Annexes K, L, M
	ISO 10542-1:2012/Cor 1:2013
34.	Technical systems and aids for disabled or handicapped persons — Wheelchair tiedown and occupant-restraint systems — Part 1: Requirements and test methods for all systems — Technical Corrigendum 1
	ISO 10865-1:2012
35.	Wheelchair containment and occupant retention systems for accessible transport vehicles designed for use by both sitting and standing passengers — Part 1: Systems for rearward-facing wheelchair-seated passengers

	ISO 10865-2:2015
36.	Wheelchair containment and occupant retention systems for accessible transport vehicles designed for use by both sitting and standing passengers — Part 2: Systems for forward-facing wheelchair-seated passengers
	ISO/TR 13570-1:2005
37.	Wheelchairs — Part 1: Guidelines for the application of the ISO 7176 series on wheelchairs
	ISO/TR 13570-2:2014
38.	Wheelchairs — Part 2: Typical values and recommended limits of dimensions, mass and manoeuvring space as determined in ISO 7176-5
	ISO 16840-1:2006
39.	Wheelchair seating — Part 1: Vocabulary, reference axis convention and measures for body segments,
	posture and postural support surfaces
	ISO 16840-2:2018
40.	Wheelchair seating — Part 2: Determination of physical and mechanical characteristics of seat cushions intended to manage tissue integrity
	ISO 16840-3:2022
41.	Wheelchair seating — Part 3: Determination of static, impact, and repetitive load strengths for
	postural support devices
	ISO 16840-4:2009
42.	Wheelchair seating — Part 4: Seating systems for use in motor vehicles
	ISO 16840-6:2015
43.	
43.	Wheelchair seating — Part 6: Simulated use and determination of the changes in properties of seat cushions

	ISO/TR 16840-9:2015
44.	Wheelchair seating — Part 9: Clinical interface pressure mapping guidelines for seating
	ISO 16840-10:2021
45.	Wheelchair seating — Part 10: Resistance to ignition of postural support devices — Requirements and test method
	ISO 16840-11:2022
46.	Wheelchair seating — Part 11: Determination of dissipation characteristics of sensible perspiration into seat cushions
	ISO 16840-12:2021
47.	Wheelchair seating — Part 12: Envelopment and immersion characterization of seat cushions using a dual semispherical indenter
	ISO 16840-13:2021
48.	Wheelchair seating — Part 13: Determination of the lateral stability property of a seat cushion
	ISO/TS 16840-14:2023
49.	Wheelchair seating — Part 14: Concepts related to managing external forces to maintain tissue integrity
	ISO 9999:2022
50.	Assistive products — Classification and terminology
	ISO 8669-1:1988
51.	Urine collection bags — Part 1: Vocabulary
	ISO 8670-1:1988
52.	Ostomy collection bags — Part 1: Vocabulary

	ISO 8670-2:1996
53.	Ostomy collection bags — Part 2: Requirements and test methods
	ISO 11948-1:1996
54.	Urine-absorbing aids — Part 1: Whole-product testing
	ISO 12505-1:2014
55.	Skin barrier for ostomy aids — Test methods — Part 1: Size, surface pH and water-absorbency
	ISO 12505-2:2016
56.	Skin barrier for ostomy aids — Test methods — Part 2: Wet integrity and adhesive strength
	ISO 15621:2017
57.	Absorbent incontinence aids for urine and/or faeces — General guidelines on evaluation
	ISO 16021:2000
58.	Urine-absorbing aids — Basic principles for evaluation of single-use adult-incontinence-absorbing aids from the perspective of users and caregivers
	ISO 17190-1:2020
59.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 1: Test method for determination of pH
	ISO 17190-2:2021
60.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 2: Test method for determination of the amount of residual acrylate monomers
	ISO 17190-3:2020
61.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 3: Test method for determination of the particle size distribution by sieve fractionation

	ISO 17190-4:2020
62.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 4: Test method for estimation of the moisture content as weight loss upon heating
	ISO 17190-5:2020
63.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 5: Test method for determination of the free swell capacity in saline by gravimetric measurement
	ISO 17190-6:2020
64.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 6: Test method for determination of the fluid retention capacity in saline solution by gravimetric measurement following centrifugation
	ISO 17190-7:2020
65.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 7: Test method for gravimetric determination of absorption against pressure
	ISO 17190-8:2020
66.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 8: Test method for determination of the permeability dependent absorption under pressure of saline solution by gravimetric measurement
	ISO 17190-9:2020
67.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 9: Test method for gravimetric determination of flow rate and bulk density
	ISO 17190-10:2020
68.	Urine-absorbing aids for incontinence — Polyacrylate superabsorbent powders — Part 10: Test method for determination of extractable polymer content by potentiometric titration
	ISO 17190-11:2001
69.	Urine-absorbing aids for incontinence — Test methods for characterizing polymer-based absorbent materials — Part 11: Determination of content of respirable particles

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70.	ISO 17191:2004 Urine-absorbing aids for incontinence — Measurement of airborne respirable polyacrylate superabsorbent materials — Determination of dust in collection cassettes by sodium atomic absorption spectrometry
71.	ISO 22748:2021 Absorbent incontinence products for urine and/or faeces — Product type names and illustrations2466
72.	ISO 24669:2021 Water-absorbent polyacrylate in urine absorbing products — Requirements
73.	ISO 17049:2013 Accessible design — Application of braille on signage, equipment and appliances
74.	ISO 17069:2020 Accessible design — Consideration and assistive products for accessible meeting
75.	ISO 19029:2016 Accessible design — Auditory guiding signals in public facilities

	Medical Face Masks
1.	EN 14683:2019+AC: 2019 Medical face masks Requirements and test methods
2.	ASTM F2100 – 19e1 Standard specification for performance of materials used in medical face masks
3.	GSO ISO 22609:2009 Clothing for protection against infectious agents - Medical face masks - Test method for resistance against penetration by synthetic blood (fixed volume, horizontally projected)
4.	ASTM F2101 – 14 Standard test method for evaluating the Bacterial Filtration Efficiency (BFE) of medical face mask materials, using a biological aerosol of staphylococcus aureus
5.	ASTM F2299 ASTM F2299/F2299M – 03(2017) Standard Test Method for Determining the Initial Efficiency of Materials Used in Medical Face Masks to Penetration by Particulates Using Latex Spheres
6.	EN 149:2001+A1:2009 Respiratory protective devices - Filtering half masks to protect against particles - Requirements, testing and marking
7.	NFPA 702 Standard for Classification of the Flammability of Wearing Apparel
8.	ASTM F1862/F1862M – 17 Standard test method for resistance of medical face masks to penetration by synthetic blood (horizontal projection of fixed volume at a known velocity)
9.	EN 143:2007 RESPIRATORY PROTECTIVE DEVICES - PARTICLE FILTERS - REQUIREMENTS, TESTING, MARKING

10.	ASTM F2100 – 19 Standard Specification For Performance Of Materials Used In Medical Face Masks
11.	16 CFR Part 1610—STANDARD FOR THE FLAMMABILITY OF CLOTHING TEXTILES



Personal Protective Equipment for Medical Purpose	
1.	ISO 4007:2018 Personal protective equipment — Eye and face protection — Vocabulary
2.	ISO 4849:1981 Personal eye-protectors — Specifications
3.	EN 166/2002 Personal eye protection
4.	OSHA 1910.133 Eye and face protection

	Complementary and Alternative Medicine	
1.	ISO 5227:2022 Traditional Chinese medicine — Safety controls for cupping devices	
2.	ISO 19611:2017 Traditional Chinese medicine — Air extraction cupping device	
3.	ISO 22213:2020 Traditional Chinese medicine — Glass cupping device	
4.	ISO 17218:2014 Sterile acupuncture needles for single use	
5.	ISO 18746:2016 Traditional Chinese medicine — Sterile intradermal acupuncture needles for single use	
6.	ISO 22236:2020 Traditional Chinese medicine —Thread-embedding acupuncture needle for single use	
7.	ISO 20308:2017 Traditional Chinese medicine — Gua Sha instruments	

Contraception						
1.	ISO 16037:2002/Amd 1:2011 Rubber condoms for clinical trials — Measurement of physical properties — Amendment 1					
2.	ISO 19671:2018 Additional lubricants for male natural rubber latex condoms — Effect on condom strength					
3.	ISO/TR 19969:2018 Guidance on sample handling for determination of bursting volume and pressure, and testing for freedom from holes for male condom					
4.	ISO/TR 24484:2023 Female condoms — Use of ISO 25841 and the quality management of female condoms					
5.	ISO 25841:2017 Female condoms — Requirements and test methods					
6.	ISO 25841:2017/Amd 1:2020 Female condoms — Requirements and test methods — Amendment 1					
7.	ISO 29942:2011 Prophylactic dams — Requirements and test methods					

Other						
1.	EN 60645-4:1995 Audiometers - Part 4: Equipment for extended high-frequency audiometry					
2.	ISO 14698-1:2003 Cleanrooms and associated controlled environments biocontamination control part 1: general principles and methods					
3.	ISO 14698-2:2003 + COR 1:2004 Cleanrooms and associated controlled environments biocontamination control part 2: evaluation and interpretation of biocontamination data					
4.	ISO 14644-1:1999 Cleanrooms and associated controlled environments — Part 1: Classification of air cleanliness by particle concentration					
5.	ISO 14644-2:2015 Cleanrooms and associated controlled environments — Part 2: Monitoring to provide evidence of cleanroom performance related to air cleanliness by particle concentration					
6.	ISO 14644-3:2019 Cleanrooms and associated controlled environments — Part 3: Test methods					
7.	ISO 14644-4:2001 Cleanrooms and associated controlled environments part 4: design, construction and start-up					
8.	ISO 14644-5:2004 Cleanrooms and associated controlled environments part 5: operations					
9.	ISO 14644-6:2007 Cleanrooms and associated controlled environments part 6: vocabulary					

	ISO 14644-7:2004
10.	Cleanrooms and associated controlled environments part 7: separative devices (clean air hoods,
	gloveboxes, isolators and mini-environments)
	ISO 14644-8:2013
11.	Cleanrooms and associated controlled environments part 8: classification of air cleanliness by
	chemical concentration (ACC)
	ISO 14644-9:2012
12.	Cleanrooms and associated controlled environments part 9: classification of surface cleanliness by
	particle concentration
	ISO 14644-10:2013
13.	Cleanrooms and associated controlled environments part 10: classification of surface cleanliness by
	chemical concentration
	IFC (0520,1090, AMD1,1000, AMD2,2012
14.	IEC 60529:1989+AMD1:1999+AMD2:2013
	Degrees of protection provided by enclosures (IP Code)
	IEC 60825-1:2014
15.	Safety of laser products - Part 1: Equipment classification and requirements
	IEC 61000-3-2:2018+AMD1:2020
1.0	
16.	Electromagnetic compatibility (EMC) - Part 3-2: Limits - Limits for harmonic current emissions (equipment input current ≤16 A per phase)
	IEC 61000-3-3:2013+AMD1:2017
17.	Electromagnetic compatibility (EMC) - Part 3-3: Limits - Limitation of voltage changes, voltage
	fluctuations and flicker in public low-voltage supply systems, for equipment with rated current ≤ 16 A per phase and not subject to conditional connection
	IEC 61000-4-2:2008
18.	Electromagnetic compatibility (EMC) - Part 4-2: Testing and measurement techniques - Electrostatic
	discharge miniumty test
18.	Electromagnetic compatibility (EMC) - Part 4-2: Testing and measurement techniques - Electrostatic discharge immunity test

19.	IEC 61000-4-3:2006+AMD1:2007+AMD2:2010 Electromagnetic compatibility (EMC) - Part 4-3: Testing and measurement techniques - Radiated, radio-frequency, electromagnetic field immunity test					
20.	IEC 61000-4-4:2012 Electromagnetic compatibility (EMC) - Part 4-4: Testing and measurement techniques - Electrical fast transient/burst immunity test					
21.	IEC 61000-4-5:2014+AMD1:2017 Electromagnetic compatibility (EMC) - Part 4-5: Testing and measurement techniques - Surge immunity test					
22.	IEC 61000-4-6:2013 Electromagnetic compatibility (EMC) - Part 4-6: Testing and measurement techniques - Immunity to conducted disturbances, induced by radio-frequency fields					
23.	IEC 61000-4-8:2009 Electromagnetic compatibility (EMC) - Part 4-8: Testing and measurement techniques - Power frequency magnetic field immunity test					
24.	IEC 61000-4-11:2020 Electromagnetic compatibility (EMC) - Part 4-11: Testing and measurement techniques - Voltage dips, short interruptions and voltage variations immunity tests for equipment with input current up to 16 A per phase					
25.	CISPR 11:2015+AMD1:2016+AMD2:2019 Industrial, scientific and medical equipment - Radio-frequency disturbance characteristics - Limits and methods of measurement					
26.	ISO 22609:2020 Clothing for protection against infectious agents medical face masks test method for resistance against penetration by synthetic blood (fixed volume, horizontally projected)					

27.	ISO 386:1977 Liquid-in-glass laboratory thermometers principles of design, construction and use				
28.	ASTM F2100 – 11(2018) Standard specification for performance of materials used in medical face masks				
29.	IEC 81001-5-1:2021 Health software and health IT systems safety, effectiveness and security				
30.	ISO 11117:2019 Gas cylinders. Valve protection caps and guards. Design, construction and tests				
31.	ISO 22882:2016 Castors and wheels — Requirements for castors for hospital beds				
32.	ISO 22610:2018 Surgical drapes, gowns and clean air suits, used as medical devices, for patients, clinical staff and equipment — Test method to determine the resistance to wet bacterial penetration				
33.	ASTM F1671/F1671M-22 Standard Test Method for Resistance of Materials Used in Protective Clothing to Penetration by Blood-Borne Pathogens Using Phi-X174 Bacteriophage Penetration as a Test System				
34.	ASTM F1980-21 Standard Guide for Accelerated Aging of Sterile Barrier Systems and Medical Devices				
35.	ISO 10282:2023 Single-use sterile rubber surgical gloves Specification				
36.	ISO 11193-1:2020 Single-use medical examination gloves Part 1: Specification for gloves made from rubber latex or rubber solution				

37. ISO 11193-2:2006 Single-use medical examination gloves Part 2: Specification for gloves made from poly(viny					
38.	EN 455-1:2020+A1:2022 Medical gloves for single use - Part 1: Requirements and testing for freedom from holes				
39.	EN 455-2:2015 Medical gloves for single use - Part 2: Requirements and testing for physical properties				
40.	EN 455-3:2023 Medical gloves for single use - Part 3: Requirements and testing for biological evaluation				
41.	EN 455-4:2009 Medical gloves for single use - Part 4: Requirements and testing for shelf life determination				
42.	ISO 7886-1:2017 Sterile hypodermic syringes for single use Part 1: Syringes for manual use				
43.	ISO 7886-2:2020 Sterile hypodermic syringes for single use Part 2: Syringes for use with power-driven syringe pumps				
44.	ISO 7886-3:2020 Sterile hypodermic syringes for single use Part 3: Auto-disabled syringes for fixed-dose immunization				
45.	ISO 7886-4:2018 Sterile hypodermic syringes for single use Part 4: Syringes with re-use prevention feature				

Annex (1): Changing to Previous Documents

Number and date of previous version		Descriptions
		Update and merge the following documents:
MDS-G44	V1.0	 SFDA Recognized Standards (Supporting Medical Device Premarket
2019/12/16		Submissions)
MDS-G46	V3.0	- Guidance on Requirements for Medical Masks - Recognized Standards
2020/06/17		
MDS - G47	V1.0	- Guidance on Requirements for Ventilators, Ventilator Tubing Connectors, and
2020/04/20		Ventilator Accessories – Rec <mark>ognized S</mark> tandards.
MDS - G004	V1.1	- Guidance for Requirements of Surgical and Medical Examination Gloves -
2022/12/14		Recognized Standards
MDS - G005	V1.1	- Guidance for Requirements of sterile single-use hypodermic syringes -
2022/12/14		Recognized Standards
MDS - G006	V1.1	- Guidance for Requirements of blood glucose metering devices and strips for
2022/12/14		home use - Recognized Standards