

Status: 2014-September			Europe					
Document Reference	Publication	Status N-Standard, N-E - Draft, VN-E predraft,	English Title	PLEASE FILL IN: IF YES = Y FOR NO = N FOR Partial = P	PLEASE FILL IN: IF YES = Y FOR NO = N FOR Partial = P	IF "Y" OR "P" PLEASE ADD THE NATIONAL/REGI ONAL REFERENCE NO.	PLEASE FILL IN:	PLEASE FILL IN:
Document Reference	Publication	Status N-Standard, N-E - Draft, VN-E predraft,	English Title	Recognised ? Y-fully, P-partial,N-NO	Mandatory ? Y-fully, P-partial,N-NO	National Reference	Publication date of the national standard	Recognition Number, if available
IEC 60118-0	1983	N	Measurement of electroacoustical characteristics					
IEC 60118-0 AMD 1	1994-01	N	Hearing aids; part_0: measurement of electroacoustical characteristics; amendment_1					
IEC 60118-1	1995-04	N	Hearing aids_- Part_1: Hearing aids with induction pick-up coil input					
IEC 60118-1 AMD 1	1998-07	N	Hearing aids_- Part_1: Hearing aids with induction pick-up coil input; Amendment_1					
IEC 60118-1 Edition 3.1	1999-01	N	Hearing aids_- Part_1: Hearing aids with induction pick-up coil input					
IEC 60118-12	1996-09	N	Hearing aids_- Part_12: Dimensions of electrical connector systems					
IEC 60118-13	2011-04	N	Electroacoustics_- Hearing aids_- Part_13: Electromagnetic compatibility (EMC)					
IEC 60118-14	1998-02	N	Hearing aids_- Part_14: Specification of a digital interface device					
IEC 60118-15	2012-02	N	Electroacoustics_- Hearing aids_- Part_15: Methods for characterising signal processing in hearing aids with a speech-like signal					
IEC 60118-2	1983	N	Hearing aids. Part 2 : Hearing aids with automatic gain control circuits					
IEC 60118-2 AMD 1	1993-02	N	Hearing aids; part_2: hearing aids with automatic gain control circuits; amendment_1					
IEC 60118-2 AMD 2	1997-05	N	Hearing aids_- Part_2: Hearing aids with automatic gain control circuits; Amendment_2					
IEC 60118-4	2006-10	N	Electroacoustics_- Hearing aids_- Part_4: Induction loop systems for hearing aid purposes_- Magnetic field strength					

IEC 60118-5	1983	N	Hearing aids. Part 5 : Nipples for insert earphones					
IEC 60118-6	1999-06	N	Hearing aids_- Part_6: Characteristics of electrical input circuits for hearing aids					
IEC 60118-7	2005-10	N	Electroacoustics_- Hearing aids_- Part_7: Measurement of performance characteristics of hearing aids for production, supply and delivery quality assurance purposes					
IEC 60118-8	2005-10	N	Electroacoustics_- Hearing aids_- Part_8: Methods of measurement of performance characteristics of hearing aids under simulated in situ working conditions					
IEC 60118-9	1985	N	Hearing aids. Part 9 : Methods of measurement of characteristics of hearing aids with bone vibrator output					
IEC 60318-4	2010-01	N	Electroacoustics_- Simulators of human head and ear_- Part_4: Occluded-ear simulator for the measurement of earphones coupled to the ear by means of ear inserts					
IEC 60335-2-52	2005-10	N	Household and similar electrical appliances_- Safety_- Part_2-52: Particular requirements for oral hygiene appliances					
IEC 60335-2-52 AMD 1	2008-04	N	Household and similar electrical appliances_- Safety_- Part_2-52: Particular requirements for oral hygiene appliances; Amendment_1					
IEC 60335-2-52 Edition 3	2008-07	N	Household and similar electrical appliances_- Safety_- Part_2-52: Particular requirements for oral hygiene appliances					
IEC 60336	2005-04	N	Medical electrical equipment_- X-ray tube assemblies for medical diagnosis_- Characteristics of focal spots					
IEC 60336 Corrigendum	2006-05	N	Medical electrical equipment_- X-ray tube assemblies for medical diagnosis_- Characteristics of focal spots; Corrigendum_1	N				
IEC 60522	2003-12	N	Determination of the permanent filtration of X-ray tube assemblies	N				
IEC 60526	1978	N	High-voltage cable plug and socket connections for medical X-ray equipment	Y				
IEC 60526 Corrigendum	2010-04	N	High-voltage cable plug and socket connections for medical X-ray equipment	N				
IEC 60580	2003-09	N	Medical electrical equipment_- Dose area product meters	N				
IEC 60601-1	2005-12	N	Medical electrical equipment_- Part_1: General requirements for basic safety and essential performance	Y				

IEC 60601-1 Corrigend	2006-12	N	Medical electrical equipment_ - Part_1: General requirements for basic safety and essential performance; Corrigendum_1	Y				
IEC 60601-1 Corrigend	2007-12	N	Medical electrical equipment_ - Part_1: General requirements for basic safety and essential performance; Corrigendum_2	Y				
IEC 60601-1 Interpreta	2008-04	N	Medical electrical equipment_ - Part_1: General requirements for basic safety and essential performance	Y				
IEC 60601-1 Interpreta	2009-01	N	Medical electrical equipment_ - Part_1: General requirements for basic safety and essential performance_ - Interpretation sheet_2	Y				
IEC 60601-1-1	2000-12	N	Medical electrical equipment_ - Part_1-1: General requirements for safety; Collateral standard: Safety requirements for medical electrical systems	Y				
IEC 60601-1-10	2007-11	N	Medical electrical equipment_ - Part_1-10: General requirements for basic safety and essential performance_ - Collateral Standard: Requirements for the development of physiologic closed-loop controllers	Y				
IEC 60601-1-11	2010-04	N	Medical electrical equipment_ - Part_1-11: General requirements for basic safety and essential performance_ - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	Y				
IEC 60601-1-11 Corrig	2011-04	N	Medical electrical equipment_ - Part_1-11: General requirements for basic safety and essential performance_ - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	Y				
IEC 60601-1-11 Techn	2011-04	N	Medical electrical equipment_ - Part_1-11: General requirements for basic safety and essential performance_ - Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment; Technical Corrigendum_1	Y				

IEC 60601-1-2	2007-03	N	Medical electrical equipment_ - Part_1-2: General requirements for basic safety and essential performance_ - Collateral standard: Electromagnetic compatibility_ - Requirements and tests	Y				
IEC 60601-1-2 Interpretation	2010-03	N	Medical electrical equipment_ - Part_1-2: General requirements for basic safety and essential performance_ - Collateral standard: Electromagnetic compatibility_ - Requirements and tests	Y				
IEC 60601-1-3	2008-01	N	Medical electrical equipment_ - Part_1-3: General requirements for basic safety and essential performance_ - Collateral standard: Radiation protection in diagnostic X-ray equipment	Y				
IEC 60601-1-4	1996-05	N	Medical electrical equipment_ - Part_1: General requirements for safety_ - 4_ - Collateral standard: Programmable electrical medical systems	Y				
IEC 60601-1-4 AMD 1	1999-10	N	Medical electrical equipment_ - Part_1-4: General requirements for safety_ - Collateral standard: Programmable electrical medical systems; Amendment_1	Y				
IEC 60601-1-4 Edition 1.	2000-04	N	Medical electrical equipment_ - Part_1-4: General requirements for safety_ - Collateral standard: Programmable electrical medical systems	Y				
IEC 60601-1-6	2010-01	N	Medical electrical equipment_ - General requirements for basic safety and essential performance_ - Collateral Standard: Usability	Y				
IEC 60601-1-8	2006-10	N	Medical electrical equipment_ - Part_1-8: General requirements for basic safety and essential performance_ - Collateral Standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	Y				
IEC 60601-1-9	2007-07	N	Medical electrical equipment_ - Part_1-9: General requirements for basic safety and essential performance_ - Collateral Standard: Requirements for environmentally conscious design	N				

IEC 60601-2-1	2009-10	N	Medical electrical equipment_ - Part_2-1: Particular requirements for the basic safety and essential performance of electron accelerators in the range 1_MeV to 50_MeV	N				
IEC 60601-2-10	1987	N	Medical electrical equipment; part_2: particular requirements for the safety of nerve and muscle stimulators	Y				
IEC 60601-2-10 AMD	2001-09	N	Medical electrical equipment_ - Part_2-10: Particular requirements for the safety of nerve and muscle stimulators; Amendment_1	Y				
IEC 60601-2-10 AMD	2002-02	N	Medical electrical equipment_ - Part_2-10: Particular requirements for the safety of nerve and muscle stimulators; Amendment_1	Y				
IEC 60601-2-11	1997-08	N	Medical electrical equipment_ - Part_2: Particular requirements for the safety of gamma beam therapy equipment	Y				
IEC 60601-2-11 AMD	2004-07	N	Amendment_1_ - Medical electrical equipment_ - Part_2-11: Particular requirements for the safety of gamma beam therapy equipment	Y				
IEC 60601-2-13	2003-05	N	Medical electrical equipment_ - Part_2-13: Particular requirements for the safety and essential performance of anaesthetic systems	Y				
IEC 60601-2-13 AMD	2006-05	N	Medical electrical equipment_ - Part_2-13: Particular requirements for the safety and essential performance of anaesthetic systems; Amendment_1	Y				
IEC 60601-2-13 Edition	2009-08	N	Medical electrical equipment_ - Part_2-13: Particular requirements for the safety of anaesthetic systems	Y				
IEC 60601-2-16	2008-04	N	Medical electrical equipment_ - Part_2-16: Particular requirements for basic safety and essential performance of haemodialysis, haemodiafiltration and haemofiltration equipment	N				
IEC 60601-2-16 Corrig	2008-10	N	Medical electrical equipment_ - Part_2-16: Particular requirements for basic safety and essential performance of haemodialysis, haemodiafiltration and haemofiltration equipment	N				
IEC 60601-2-17	2005-09	N	Medical electrical equipment_ - Part_2-17: Particular requirements for the safety of automatically-controlled brachytherapy afterloading equipment	N				

IEC 60601-2-18	2009-08	N	Medical electrical equipment_ - Part_2-18: Particular requirements for basic safety and essential performance of endoscopic equipment	Y				
IEC 60601-2-19	2009-02	N	Medical electrical equipment_ - Part_2-19: Particular requirements for the basic safety and essential performance of infant incubators	N				
IEC 60601-2-19 Corrig	2012-02	N	Medical electrical equipment_ - Part_2-19: Particular requirements for the basic safety and essential performance of infant incubators; Corrigendum_1	N				
IEC 60601-2-2	2009-02	N	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	N				
IEC 60601-2-20	2009-02	N	Medical electrical equipment_ - Part_2-20: Particular requirements for the basic safety and essential performance of infant transport incubators	Y				
IEC 60601-2-20 Corrig	2012-02	N	Medical electrical equipment_ - Part_2-20: Particular requirements for the basic safety and essential performance of infant transport incubators; Corrigendum_1	N				
IEC 60601-2-21	2009-02	N	Medical electrical equipment_ - Part_2-21: Particular requirements for the basic safety and essential performance of infant radiant warmers	N				
IEC 60601-2-22	2007-05	N	Medical electrical equipment_ - Part_2-22: Particular requirements for basic safety and essential performance of surgical, cosmetic, therapeutic and diagnostic laser equipment	N				
IEC 60601-2-23	2011-02	N	Medical electrical equipment_ - Part_2-23: Particular requirements for the basic safety and essential performance of transcutaneous partial pressure monitoring equipment	N				
IEC 60601-2-24	1998-02	N	Medical electrical equipment_ - Part_2-24: Particular requirements for the safety of infusion pumps and controllers	N				

IEC 60601-2-25	2011-10	N	Medical electrical equipment_ - Part_2-25: Particular requirements for basic safety and essential performance of electrocardiographs	Y				
IEC 60601-2-26	2003-12	N	Medical electrical equipment_ - Part_2-26: Particular requirements for the safety of electroencephalographs	N				
IEC 60601-2-27	2011-03	N	Medical electrical equipment_ - Part_2-27: Particular requirements for the basic safety and essential performance of electrocardiographic monitoring equipment	Y				
IEC 60601-2-28	2010-03	N	Medical electrical equipment_ - Part_2-28: Particular requirements for basic safety and essential performance of X-ray tube assemblies for medical diagnosis	N				
IEC 60601-2-29	2008-06	N	Medical electrical equipment_ - Part_2-29: Particular requirements for the basic safety and essential performance of radiotherapy simulators	Y				
IEC 60601-2-3	1991-06	N	Medical electrical equipment; part_2: particular requirements for the safety of short-wave therapy equipment	Y				
IEC 60601-2-3 AMD 1	1998-09	N	Medical electrical equipment_ - Part_2: Particular requirements for the safety of short-wave therapy equipment; Amendment_1	Y				
IEC 60601-2-31	2008-03	N	Medical electrical equipment_ - Part_2-31: Particular requirements for basic safety and essential performance of external cardiac pacemakers with internal power source	Y				
IEC 60601-2-31 AMD	2011-06	N	Medical electrical equipment_ - Part_2-31: Particular requirements for basic safety and essential performance of external cardiac pacemakers with internal power source	N				
IEC 60601-2-31 Edition	2011-09	N	Medical electrical equipment_ - Part_2-31: Particular requirements for basic safety and essential performance of external cardiac pacemakers with internal power source	N				
IEC 60601-2-32	1994-03	N	Medical electrical equipment; part_2: particular requirements for the safety of X-ray equipment	N				

IEC 60601-2-33	2010-03	N	Medical electrical equipment_ - Part_2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis	Y				
IEC 60601-2-33 Corriger	2012-03	N	Medical electrical equipment_ - Part_2-33: Particular requirements for the basic safety and essential performance of magnetic resonance equipment for medical diagnosis	Y				
IEC 60601-2-34	2011-05	N	Medical electrical equipment_ - Part_2-34: Particular requirements for the basic safety and essential performance of invasive blood pressure monitoring equipment	N				
IEC 60601-2-36	1997-03	N	Medical electrical equipment_ - Part_2: Particular requirements for the safety of equipment for extracorporeally induced lithotripsy	N				
IEC 60601-2-37	2007-08	N	Medical electrical equipment_ - Part_2-37: Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment	Y				
IEC 60601-2-39	2007-11	N	Medical electrical equipment_ - Part_2-39: Particular requirements for basic safety and essential performance of peritoneal dialysis equipment	Y				
IEC 60601-2-4	2010-12	N	Medical electrical equipment_ - Part_2-4: Particular requirements for basic safety and essential performance of cardiac defibrillators	Y				
IEC 60601-2-40	1998-02	N	Medical electrical equipment_ - Part_2-40: Particular requirements for the safety of electromyographs and evoked response equipment	N				
IEC 60601-2-41	2009-08	N	Medical electrical equipment_ - Part_2-41: Particular requirements for basic safety and essential performance of surgical luminaires and luminaires for diagnosis	Y				
IEC 60601-2-43	2010-03	N	Medical electrical equipment_ - Part_2-43: Particular requirements for basic safety and essential performance of X-ray equipment for interventional procedures	N				

IEC 60601-2-44	2009-02	N	Medical electrical equipment_ - Part_2-44: Particular requirements for the basic safety and essential performance of X-ray equipment for computed tomography	Y				
IEC 60601-2-44 Corrig	2010-05	N	Medical electrical equipment_ - Part_2-44: Particular requirements for the basic safety and essential performance of X-ray equipment for computed tomography	Y				
IEC 60601-2-45	2011-02	N	Medical electrical equipment_ - Part_2-45: Particular requirements for the basic safety and essential performance of mammographic X-ray equipment and mammographic stereotactic devices	Y				
IEC 60601-2-46	2010-12	N	Medical electrical equipment_ - Part_2-46: Particular requirements for the basic safety and essential performance of operating tables	N				
IEC 60601-2-47	2012-02	N	Medical electrical equipment_ - Part_2-47: Particular requirements for the basic safety and essential performance of ambulatory electrocardiographic systems	N				
IEC 60601-2-49	2011-02	N	Medical electrical equipment_ - Part_2-49: Particular requirements for the basic safety and essential performance of multifunction patient monitoring equipment	N				
IEC 60601-2-5	2009-07	N	Medical electrical equipment_ - Part_2-5: Particular requirements for basic safety and essential performance of ultrasonic physiotherapy equipment	N				
IEC 60601-2-50	2009-03	N	Medical electrical equipment_ - Part_2-50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment	N				
IEC 60601-2-50 Corrig	2010-08	N	Medical electrical equipment_ - Part_2-50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment	Y				
IEC 60601-2-52	2009-12	N	Medical electrical equipment_ - Part_2-52: Particular requirements for the basic safety and essential performance of medical beds	Y				

IEC 60601-2-52 Corrig	2010-09	N	Medical electrical equipment_ - Part_2-52: Particular requirements for the basic safety and essential performance of medical beds	Y				
IEC 60601-2-52 Techn	2010-09	N	Medical electrical equipment_ - Part_2-52: Particular requirements for the basic safety and essential performance of medical beds; Technical Corrigendum_1	Y				
IEC 60601-2-54	2009-06	N	IEC_60601-2-54, Ed._1: Medical electrical equipment_ - Part_2-54: Particular requirements for the basic safety and essential performance of X-ray equipment for radiography and radioscopy	Y				
IEC 60601-2-54 Corrig	2010-03	N	Medical electrical equipment_ - Part_2-54: Particular requirements for the basic safety and essential performance of X-ray equipment for radiography and radioscopy	Y				
IEC 60601-2-54 Corrig	2011-06	N	Medical electrical equipment_ - Part_2-54: Particular requirements for the basic safety and essential performance of X-ray equipment for radiography and radioscopy	Y				
IEC 60601-2-57	2011-01	N	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	Y				
IEC 60601-2-6	1984	N	Medical electrical equipment. Part 2: Particular requirements for the safety of microwave therapy equipment	N				
IEC 60601-2-7	1998-02	N	Medical electrical equipment_ - Part_2-7: Particular requirements for the safety of high-voltage generators of diagnostic X-ray generators	N				
IEC 60601-2-8	2010-11	N	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	Y				

IEC 60601-2-8 AMD 1	1997-08	N	Medical electrical equipment_ - Part_2: Particular requirements for the safety of therapeutic X-ray equipment in the range 10_kV to 1_MV; Amendment_1	N				
IEC 60601-2-8 Edition	1999-04	N	Medical electrical equipment_ - Part_2-8: Particular requirements for the safety of therapeutic X-ray equipment operating in the range 10_kV to 1_MV	N				
IEC 60601-3-1	1996-07	N	Medical electrical equipment_ - Part_3-1: Essential performance requirements for transcutaneous oxygen and carbon dioxide partial pressure monitoring equipment	N				
IEC 60613	2010-01	N	Electrical and loading characteristics of X-ray tube assemblies for medical diagnosis	N				
IEC 60627	2001-08	N	Diagnostic X-ray imaging equipment_ - Characteristics of general purpose and mammographic anti-scatter grids	Y				
IEC 60645-1	2012-02	N	Electroacoustics_ - Audiometric equipment_ - Part_1: Equipment for pure-tone audiometry	N				
IEC 60645-2	1993-11	N	Audiometers; part_2: equipment for speech audiometry	Y				
IEC 60645-3	2007-03	N	Electroacoustics_ - Audiometric equipment_ - Part_3: Test signals of short duration	Y				
IEC 60645-5	2004-11	N	Electroacoustics_ - Audiometric equipment_ - Part_5: Instruments for the measurement of aural acoustic impedance/admittance	N				
IEC 60645-6	2009-04	N	Electroacoustics_ - Audiometric equipment_ - Part_6: Instruments for the measurement of otoacoustic emissions	N				
IEC 60645-7	2009-04	N	Electroacoustics_ - Audiometric equipment_ - Part_7: Instruments for the measurement of auditory brainstem responses	N				
IEC 60789	2005-10	N	Medical electrical equipment_ - Characteristics and test conditions of radionuclide imaging devices_ - Anger type gamma cameras	N				
IEC 60789 Corrigendum	2009-10	N	Medical electrical equipment_ - Characteristics and test conditions of radionuclide imaging devices_ - Anger type gamma cameras; Corrigendum_1	N				

IEC 60806	1984	N	Determination of the maximum symmetrical radiation field from a rotating anode X-ray tube for medical diagnosis	N				
IEC 60976	2007-10	N	Medical electrical equipment_ - Medical electron accelerators_ - Functional performance characteristics	N				
IEC 61010-2-040	2005-04	N	Safety requirements for electrical equipment for measurement, control and laboratory use_ - Part_2-040: Particular requirements for sterilizers and washer-disinfectors used to treat medical materials	N				
IEC 61010-2-101	2002-01	N	Safety requirements for electrical equipment for measurement, control and laboratory use_ - Part_2-101: Particular requirements for in vitro diagnostic (IVD) medical equipment	Y				
IEC 61157	2007-08	N	Standard means for the reporting of the acoustic output of medical diagnostic ultrasonic equipment	N				
IEC 61157 Corrigendum	2008-08	N	Standard means for the reporting of the acoustic output of medical diagnostic ultrasonic equipment; Corrigendum_1	N				
IEC 61168	1993-12	N	Radiotherapy simulators; functional performance characteristics	N				
IEC 61205	1993-12	N	Ultrasonics; dental descaler systems; measurement and declaration of the output characteristics					
IEC 61217	2011-12	N	Radiotherapy equipment coordinates, movements and scales	N				
IEC 61223-2-6	2006-11	N	Evaluation and routine testing in medical imaging departments_ - Part_2-6: Constancy tests_ - Imaging performance of computed tomography X-ray equipment	N				
IEC 61223-3-2	2007-07	N	Evaluation and routine testing in medical imaging departments_ - Part_3-2: Acceptance tests_ - Imaging performance of mammographic X-ray equipment	N				
IEC 61223-3-4	2000-03	N	Evaluation and routine testing in medical imaging departments_ - Part_3-4: Acceptance tests_ - Imaging performance of dental X-ray equipment	N				

IEC 61223-3-5	2004-08	N	Evaluation and routine testing in medical imaging departments_- Part_3-5: Acceptance tests_- Imaging performance of computed tomography X-ray equipment	N				
IEC 61223-3-5 Corrige	2006-03	N	Evaluation and routine testing in medical imaging departments_- Part_3-5: Acceptance tests_- Imaging performance of computed tomography X-ray equipment; Corrigendum_1	N				
IEC 61252 Edition 1.1	2002-03	N	Electroacoustics_- Specifications for personal sound exposure meters	N				
IEC 61262-1	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_1: Determination of the entrance field size	N				
IEC 61262-2	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_2: Determination of the conversion factor	N				
IEC 61262-3	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_3: Determination of the luminance distribution and luminance non-uniformity	N				
IEC 61262-4	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_4: Determination of the image distortion	N				
IEC 61262-5	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_5: Determination of the detective quantum efficiency	N				
IEC 61262-6	1994-07	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part_6: Determination of the contrast ratio and veiling glare index	N				
IEC 61262-7	1995-09	N	Medical electrical equipment_- Characteristics of electro-optical X-ray image intensifiers_- Part-7: Determination of the modulation transfer function	N				

IEC 61266	1994-12	N	Ultrasonics_- Hand-held probe Doppler foetal heartbeat detectors_- Performance requirements and methods of measurement and reporting	N				
IEC 61267	2005-11	N	Medical diagnostic X-ray equipment_- Radiation conditions for use in the determination of characteristics	N				
IEC 61303	1994-09	N	Medical electrical equipment_- Radionuclide calibrators_- Particular methods for describing performance	N				
IEC 61326-2-6	2005-12	N	Electrical equipment for measurement, control and laboratory use, control and laboratory use_- EMC requirements_- Part_2-6: Particular requirements_- In-vitro diagnostic (IVD) medical equipment	Y				
IEC 61326-2-6 Corrigé	2007-09	N	Electrical equipment for measurement, control and laboratory use, control and laboratory use_- EMC requirements_- Part_2-6: Particular requirements_- In-vitro diagnostic (IVD) medical equipment; Corrigendum_1	N				
IEC 61331-1	1994-10	N	Protective devices against diagnostic medical X-radiation_- Part_1: Determination of attenuation properties of materials	N				
IEC 61331-2	1994-10	N	Protective devices against diagnostic medical X-radiation_- Part_2: Protective glass plates	N				
IEC 61331-3	1998-11	N	Protective devices against diagnostic medical X-radiation_- Part_3: Protective clothing and protective devices for gonads	N				
IEC 61391-1	2006-07	N	Ultrasonics_- Pulse echo scanners_- Part_1: Techniques for calibrating spatial measurement systems and measurement of system point-spread function response	N				
IEC 61391-2	2010-01	N	Ultrasonics_- Pulse-echo scanners_- Part_2: Measurement of maximum depth of penetration and local dynamic range	N				
IEC 61669	2001-01	N	Electroacoustics_- Equipment for the measurement of real-ear acoustical characteristics of hearing aids					
IEC 61674 AMD 1	2002-06	N	Medical electrical equipment_- Dosimeters with ionization chambers and/or semi-conductor detectors as used in X-ray diagnostic imaging; Amendment_1	N				

IEC 61675-1	1998-02	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_1: Positron emission tomographs	N				
IEC 61675-1 AMD 1	2008-04	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_1: Positron emission tomographs; Amendment_1	N				
IEC 61675-1 Edition 1.	2008-06	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_1: Positron emission tomographs	N				
IEC 61675-2	1998-01	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_2: Single photon emission computed tomographs	N				
IEC 61675-2 AMD 1	2004-12	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_2: Single photon emission computed tomographs; Amendment_1	N				
IEC 61675-2 Edition 1.	2005-02	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_2: Single photon emission computed tomographs	N				
IEC 61675-3	1998-02	N	Radionuclide imaging devices_- Characteristics and test conditions_- Part_3: Gamma camera based wholebody imaging systems	N				
IEC 61676	2002-09	N	Medical electrical equipment_- Dosimetric instruments used for non-invasive measurement of X-ray tube voltage in diagnostic radiology	Y				
IEC 61676 AMD 1	2008-11	N	Medical electrical equipment_- Dosimetric instruments used for non-invasive measurement of x-ray tube voltage in diagnostic radiology; Amendment_1	Y				
IEC 61676 Edition 1.1	2009-01	N	Medical electrical equipment_- Dosimetric instruments used for non-invasive measurement of X-ray tube voltage in diagnostic radiology	Y				
IEC 61685	2002-09	N	Medical electrical equipment_- Dosimetric instruments used for non-invasive measurement of X-ray tube voltage in diagnostic radiology					
IEC 61689	2007-08	N	Ultrasonics_- Physiotherapy systems_- Field specifications and methods of measurement in the frequency range 0,5_MHz to 5_MHz	N				
IEC 61846	1998-04	N	Ultrasonics_- Pressure pulse lithotripters_- Characteristics of fields	N				

IEC 61847	1998-01	N	Ultrasonics_ - Surgical systems_ - Measurement and declaration of the basic output characteristics	N				
IEC 62083	2009-09	N	Medical electrical equipment_ - Requirements for the safety of radiotherapy treatment planning systems	Y				
IEC 62127.1	2003-10	N	Medical electrical equipment_ - Characteristics of digital X-ray imaging devices_ - Part_1: Determination of the detective quantum efficiency					
IEC 62220-1	2003-10	N	Medical electrical equipment_ - Characteristics of digital X-ray imaging devices_ - Part_1: Determination of the detective quantum efficiency	Y				
IEC 62220-1-2	2007-06	N	Medical electrical equipment_ - Characteristics of digital X-ray imaging devices_ - Part_1-2: Determination of the detective quantum efficiency_ - Detectors used in mammography	Y				
IEC 62220-1-3	2008-06	N	Medical electrical equipment_ - Characteristics of digital X-ray imaging devices_ - Part_1-3: Determination of the detective quantum efficiency_ - Detectors used in dynamic imaging	Y				
IEC 62274	2005-05	N	Medical electrical equipment_ - Safety of radiotherapy record and verify systems	N				
IEC 62304	2006-05	N	Medical device software_ - Software life cycle processes	Y				
IEC 62353	2007-05	N	Medical electrical equipment_ - Recurrent test and test after repair of medical electrical equipment	N				
IEC 62359	2010-10	N	Ultrasonics_ - Field characterization_ - Test methods for the determination of thermal and mechanical indices related to medical diagnostic ultrasonic fields	N				
IEC 62359 Corrigendum	2011-03	N	Ultrasonics_ - Field characterization_ - Test methods for the determination of thermal and mechanical indices related to medical diagnostic ultrasonic fields	N				
IEC 62366	2007-10	N	Medical devices_ - Application of usability engineering to medical devices	Y				
IEC 62464-1	2007-01	N	Magnetic resonance equipment for medical imaging_ - Part_1: Determination of essential image quality parameters	N				

IEC 62464-2	2010-11	N	Magnetic resonance equipment for medical imaging_ - Part_2: Classification criteria for pulse sequences	N				
IEC 62489-1	2010-01	N	Electroacoustics_ - Audio-frequency induction loop systems for assisted hearing_ - Part_1: Methods of measuring and specifying the performance of system components					
IEC 62489-2	2011-01	N	Electroacoustics_ - Audio-frequency induction loop systems for assisted hearing_ - Part_2: Methods of calculating and measuring the low-frequency magnetic field emissions from the loop for assessing conformity with guidelines on limits for human exposure					
IEC 62494-1	2008-08	N	Medical electrical equipment_ - Exposure index of digital X-ray imaging systems_ - Part_1: Definition and requirements of general radiography	N				
IEC 62563-1	2009-12	N	Medical electrical equipment_ - Medical image display systems_ - Part_1: Evaluation methods	N				
IEC 80001-1	2010-10	N	Application of risk management for IT-networks incorporating medical devices_ - Part_1: Roles, responsibilities and activities	N				
IEC 80601-2-30	2009-01	N	Medical electrical equipment_ - Part_2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers	N				
IEC 80601-2-30 Corrig	2010-01	N	Medical electrical equipment_ - Part_2-30: Particular requirements for basic safety and essential performance of automated non-invasive sphygmomanometers; Corrigendum_1	N				
IEC 80601-2-35	2009-10	N	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	Y				
IEC 80601-2-35 Corrig	2012-03	N	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	N				

IEC 80601-2-58	2008-10	N	Medical electrical equipment_ - Part_2-58: Particular requirements for basic safety and essential performance of lens removal devices and vitrectomy devices for ophthalmic surgery	Y				
IEC 80601-2-59	2008-10	N	Medical electrical equipment_ - Part_2-59: Particular requirements for basic safety and essential performance of screening thermographs for human febrile temperature screening	Y				
IEC 80601-2-59 Corrig	2009-04	N	Medical electrical equipment_ - Part_2-59: Particular requirements for basic safety and essential performance of screening thermographs for human febrile temperature screening; Corrigendum_1	N				
IEC 80601-2-60	2012-02	N	Medical electrical equipment_ - Part_2-60: Particular requirements for basic safety and essential performance of dental equipment					
IEC/TR 60788	2004-02	N	Medical electrical equipment_ - Glossary of defined terms	N				
IEC/TR 60825-8	2006-12	N	Safety of laser products_ - Part_8: Guidelines for the safe use of laser beams on humans					
IEC/TR 60854	1986	N	Methods of measuring the performance of ultrasonic pulse-echo diagnostic equipment	N				
IEC/TR 60878	2003-07	N	Graphical symbols for electrical equipment in medical practice	N				
IEC/TR 60930	2008-09	N	Guidelines for administrative, medical, and nursing staff concerned with the safe use of medical electrical equipment and medical electrical systems	N				
IEC/TR 60977	2008-07	N	Medical electrical equipment_ - Medical electron accelerators_ - Guidelines for functional performance characteristics	N				
IEC/TR 61258	2008-08	N	Guidelines for the development and use of medical electrical equipment educational materials	N				
IEC/TR 61289	2011-11	N	High frequency surgical equipment_ - Operation and maintenance	N				
IEC/TR 61948-2	2001-02	N	Nuclear medicine instrumentation_ - Routine tests_ - Part_2: Scintillation cameras and single photon emission computed tomography imaging	N				
IEC/TR 61948-3	2005-07	N	Nuclear medicine instrumentation_ - Routine tests_ - Part_3: Positron emission tomographs	N				

IEC/TR 61948-4	2006-11	N	Nuclear medicine instrumentation_- Routine tests_- Part_4: Radionuclide calibrators	N				
IEC/TR 62266	2002-03	N	Medical electrical equipment_- Guidelines for implementation of DICOM in radiotherapy	N				
IEC/TR 62296	2009-01	N	Considerations of unaddressed safety aspects in the second edition of IEC_60601-1 and proposals for new requirements	N				
IEC/TR 62348	2006-05	N	Mapping between the clauses of the third edition of IEC_60601-1 and the 1988 edition as amended	N				
IEC/TR 62354	2009-10	N	General testing procedures for medical electrical equipment	N				
IEC/TR 62649	2010-04	N	Requirements for measurement standards for high intensity therapeutic ultrasound (HITU) devices	N				
IEC/TR 62678	2010-10	N	Audio, video and multimedia systems and equipment_- Activities and considerations related to accessibility and usability					
IEC/TR 80002-1	2009-09	N	Medical device software_- Part_1: Guidance on the application of ISO_14971 to medical device software	N				
IEC/TR2 61170	1993-12	N	Radiotherapy simulators; guidelines for functional performance characteristics	N				
IEC/TR2 61223-1	1993-07	N	Evaluation and routine testing in medical imaging departments; part_1: general aspects	N				
IEC/TR2 61390	1996-07	N	Ultrasonics_- Real-time pulse-echo systems_- Test procedures to determine the performance specifications	N				
IEC/TR3 60513	1994-01	N	Fundamental aspects of safety standards for medical electrical equipment	N				
IEC/TR3 61288-1	1993-10	N	Cardiac defibrillators; cardiac defibrillators-monitors; part_1: operation	N				
IEC/TR3 61288-2	1993-10	N	Cardiac defibrillators; cardiac defibrillators-monitors; part_2: maintenance	N				
IEC/TR3 61852	1998-04	N	Medical electrical equipment_- Digital imaging and communications in medicine (DICOM)_- Radiotherapy objects	N				
IEC/TR3 61859	1997-04	N	Guidelines for radiotherapy treatment rooms design	N				

ISO 10079-1	1999-08	N	Medical suction equipment_ - Part_1: Electrically powered suction equipment_ - Safety requirements	Y				
ISO 10079-2	1999-08	N	Medical suction equipment_ - Part_2: Manually powered suction equipment	Y				
ISO 10079-3	1999-08	N	Medical suction equipment_ - Part_3: Suction equipment powered from a vacuum or pressure source	Y				
ISO 10083	2006-07	N	Oxygen concentrator supply systems for use with medical gas pipeline systems	N				
ISO 10139-1	2005-02	N	Dentistry_ - Soft lining materials for removable dentures_ - Part_1: Materials for short-term use					
ISO 10139-1 Technical C	2006-03	N	Dentistry_ - Soft lining materials for removable dentures_ - Part_1: Materials for short-term use; Technical Corrigendum_1					
ISO 10139-2	2009-08	N	Dentistry_ - Soft lining materials for removable dentures_ - Part_2: Materials for long-term use					
ISO 10159	2011-12	N	Health informatics_ - Messages and communication_ - Web access reference manifest					
ISO 10271	2011-08	N	Dentistry_ - Corrosion test methods for metallic materials					
ISO 10282	2002-09	N	Single-use sterile rubber surgical gloves_ - Specification	N				
ISO 10282 Technical C	2005-06	N	Single-use sterile rubber surgical gloves_ - Specification; Technical Corrigendum_1	N				
ISO 10322-1	2006-02	N	Ophthalmic optics_ - Semi-finished spectacle lens blanks_ - Part_1: Specifications for single-vision and multifocal lens blanks	N				
ISO 10322-2	2006-02	N	Ophthalmic optics_ - Semi-finished spectacle lens blanks_ - Part_2: Specifications for progressive power lens blanks	N				
ISO 10323	1991-11	N	Dental rotary instruments; bore diameters for discs and wheels					
ISO 10328	2006-10	N	Prosthetics_ - Structural testing of lower-limb prostheses_ - Requirements and test methods	Y				
ISO 10334	1994-08	N	Implants for surgery_ - Malleable wires for use as sutures and other surgical applications	N				
ISO 10341	2009-07	N	Ophthalmic instruments_ - Refractor heads	N				
ISO 10342	2010-06	N	Ophthalmic instruments_ - Eye refractometers	N				

ISO 10343	2009-07	N	Ophthalmic instruments_ - Ophthalmometers	N				
ISO 10451	2010-06	N	Dentistry_ - Contents of technical file for dental implant systems					
ISO 10477	2004-10	N	Dentistry_ - Polymer-based crown and bridge materials					
ISO 10524-1	2006-02	N	Pressure regulators for use with medical gases_ - Part_1: Pressure regulators and pressure regulators with flow-metering devices	Y				
ISO 10524-2	2005-05	N	Pressure regulators for use with medical gases_ - Part_2: Manifold and line pressure regulators	Y				
ISO 10524-3	2005-05	N	Pressure regulators for use with medical gases_ - Part_3: Pressure regulators integrated with cylinder valves	Y				
ISO 10524-4	2008-06	N	Pressure regulators for use with medical gases_ - Part_4: Low-pressure regulators	Y				
ISO 10535	2006-12	N	Hoists for the transfer of disabled persons_ - Requirements and test methods	Y				
ISO 10542-1	2001-07	N	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 10542-2	2001-07	N	Technical systems and aids for disabled or handicapped persons_ - Wheelchair tiedown and occupant-restraint systems_ - Part_2: Four-point strap-type tiedown systems					
ISO 10542-3	2005-02	N	Technical systems and aids for disabled or handicapped persons_ - Wheelchair tiedown and occupant-restraint systems_ - Part_3: Docking-type tiedown systems					
ISO 10542-4	2004-09	N	Technical systems and aids for disabled or handicapped persons_ - Wheelchair tiedown and occupant-restraint systems_ - Part_4: Clamp-type tiedown systems					
ISO 10542-5	2004-04	N	Technical systems and aids for disabled or handicapped persons_ - Wheelchair tiedown and occupant-restraint systems_ - Part_5: Systems for specific wheelchairs					
ISO 10555-1	1995-06	N	Sterile, single-use intravascular catheters_ - Part_1: General requirements	Y				
ISO 10555-1 AMD 1	1999-07	N	Sterile, single-use intravascular catheters_ - Part_1: General requirements; Amendment_1	Y				

ISO 10555-1 AMD 2	2004-05	N	Sterile, single-use intravascular catheters_- Part_1: General requirements; Amendment_2	Y				
ISO 10555-2	1996-06	N	Sterile, single-use intravascular catheters_- Part_2: Angiographic catheters	N				
ISO 10555-2 Technical	2002-06	N	Sterile, single-use intravascular catheters_- Part_2: Angiographic catheters; Technical Corrigendum_1	N				
ISO 10555-3	1996-06	N	Sterile, single-use intravascular catheters_- Part_3: Central venous catheters	N				
ISO 10555-3 Technical	2002-06	N	Sterile, single-use intravascular catheters_- Part_3: Central venous catheters; Technical Corrigendum_1	N				
ISO 10555-4	1996-06	N	Sterile, single-use intravascular catheters_- Part_4: Balloon dilatation catheters	N				
ISO 10555-4 Technical	2002-06	N	Sterile, single-use intravascular catheters_- Part_4: Balloon dilatation catheters; Technical Corrigendum_1	N				
ISO 10555-5	1996-06	N	Sterile, single-use intravascular catheters_- Part_5: Over-needle peripheral catheters	N				
ISO 10555-5 AMD 1	1999-01	N	Sterile, single-use intravascular catheters_- Part_5: Over-needle peripheral catheters; Amendment_1	N				
ISO 10555-5 Technical	2002-06	N	Sterile, single-use intravascular catheters_- Part_5: Over-needle peripheral catheters; Technical Corrigendum_1	N				
ISO 10637	1999-08	N	Dental equipment_- High- and medium-volume suction systems					
ISO 10650-1	2004-11	N	Dentistry_- Powered polymerization activators_- Part_1: Quartz tungsten halogen lamps					
ISO 10650-2	2007-09	N	Dentistry_- Powered polymerization activators_- Part_2: Light-emitting diode (LED) lamps					
ISO 10651-2	2004-07	N	Lung ventilators for medical use_- Particular requirements for basic safety and essential performance_- Part_2: Home care ventilators for ventilator-dependent patients	Y				
ISO 10651-3	1997-01	N	Lung ventilators for medical use_- Part_3: Particular requirements for emergency and transport ventilators	N				

ISO 10651-4	2002-03	N	Lung ventilators_ - Part_4: Particular requirements for operator-powered resuscitators	Y				
ISO 10651-5	2006-02	N	Lung ventilators for medical use_ - Particular requirements for basic safety and essential performance_ - Part_5: Gas-powered emergency resuscitators	N				
ISO 10651-6	2004-07	N	Lung ventilators for medical use_ - Particular requirements for basic safety and essential performance_ - Part_6: Home-care ventilatory support devices	Y				
ISO 10685-1	2011-12	N	Ophthalmic optics_ - Spectacle frames and sunglasses electronic catalogue and identification_ - Part_1: Product identification and electronic catalogue product hierarchy	N				
ISO 10873	2010-09	N	Dentistry_ - Denture adhesives					
ISO 10936-1	2000-06	N	Optics and optical instruments_ - Operation microscopes_ - Part_1: Requirements and test methods	N				
ISO 10936-2	2010-01	N	Optics and photonics_ - Operation microscopes_ - Part_2: Light hazard from operation microscopes used in ocular surgery	N				
ISO 10938	1998-05	N	Ophthalmic instruments_ - Chart projectors	N				
ISO 10939	2007-02	N	Ophthalmic instruments_ - Slit-lamp microscopes	N				
ISO 10940	2009-08	N	Ophthalmic instruments_ - Fundus cameras	N				
ISO 10942	2006-06	N	Ophthalmic instruments_ - Direct ophthalmoscopes	N				
ISO 10943	2011-08	N	Ophthalmic instruments_ - Indirect ophthalmoscopes	N				
ISO 10944	2009-08	N	Ophthalmic instruments_ - Synoptophores	N				
ISO 10985	2009-02	N	Caps made of aluminium-plastics combinations for infusion bottles and injection vials_ - Requirements and test methods	N				
ISO 10993-1	2009-10	N	Biological evaluation of medical devices_ - Part_1: Evaluation and testing within a risk management process	Y				

ISO 10993-1 Technical	2010-06	N	Biological evaluation of medical devices_ - Part_1: Evaluation and testing within a risk management process; Technical Corrigendum_1	Y				
ISO 10993-10	2010-08	N	Biological evaluation of medical devices_ - Part_10: Tests for irritation and skin sensitization	N				
ISO 10993-11	2006-08	N	Biological evaluation of medical devices_ - Part_11: Tests for systemic toxicity	Y				
ISO 10993-12	2007-11	N	Biological evaluation of medical devices_ - Part_12: Sample preparation and reference materials	Y				
ISO 10993-13	2010-06	N	Biological evaluation of medical devices_ - Part_13: Identification and quantification of degradation products from polymeric medical devices	Y				
ISO 10993-14	2001-11	N	Biological evaluation of medical devices_ - Part_14: Identification and quantification of degradation products from ceramics	Y				
ISO 10993-15	2000-12	N	Biological evaluation of medical devices_ - Part_15: Identification and quantification of degradation products from metals and alloys	Y				
ISO 10993-16	2010-02	N	Biological evaluation of medical devices_ - Part_16: Toxicokinetic study design for degradation products and leachables	Y				
ISO 10993-17	2002-12	N	Biological evaluation of medical devices_ - Part_17: Establishment of allowable limits for leachable substances	Y				
ISO 10993-18	2005-07	N	Biological evaluation of medical devices_ - Part_18: Chemical characterization of materials	Y				
ISO 10993-2	2006-07	N	Biological evaluation of medical devices_ - Part_2: Animal welfare requirements	N				
ISO 10993-3	2003-10	N	Biological evaluation of medical devices_ - Part_3: Tests for genotoxicity, carcinogenicity and reproductive toxicity	Y				
ISO 10993-4	2002-10	N	Biological evaluation of medical devices_ - Part_4: Selection of test for interactions with blood	Y				

ISO 10993-4 AMD 1	2006-07	N	Biological evaluation of medical devices_ - Part_4: Selection of tests for interactions with blood	Y				
ISO 10993-5	2009-06	N	Biological evaluation of medical devices_ - Part_5: Tests for in vitro cytotoxicity	Y				
ISO 10993-6	2007-04	N	Biological evaluation of medical devices_ - Part_6: Tests for local effects after implantation	Y				
ISO 10993-7	2008-10	N	Biological evaluation of medical devices_ - Part_7: Ethylene oxide sterilization residuals	Y				
ISO 10993-7 Technical	2009-11	N	Biological evaluation of medical devices_ - Part_7: Ethylene oxide sterilization residuals; Technical Corrigendum_1	Y				
ISO 10993-9	2009-12	N	Biological evaluation of medical devices_ - Part_9: Framework for identification and quantification of potential degradation products	Y				
ISO 11040-1	1992-11	N	Prefilled syringes; part_1: glass cylinders for dental local anaesthetic cartridges	N				
ISO 11040-2	2011-04	N	Prefilled syringes_ - Part_2: Plunger stoppers for dental local anaesthetic cartridges	N				
ISO 11040-3	2012-01	N	Prefilled syringes_ - Part_3: Seals for dental local anaesthetic cartridges	N				
ISO 11040-4	2007-02	N	Prefilled syringes_ - Part_4: Glass barrels for injectables	N				
ISO 11040-5	2012-01	N	Prefilled syringes_ - Part_5: Plunger stoppers for injectables	N				
ISO 11070	1998-05	N	Sterile single-use intravascular catheter introducers	N				
ISO 11073-90101	2008-01	N	Health informatics_ - Point-of-care medical device communication_ - Part_90101: Analytical instruments_ - Point-of-care test	N				
ISO 11073-91064	2009-05	N	Health informatics_ - Standard communication protocol_ - Part_91064: Computer-assisted electrocardiography					
ISO 11135-1	2007-05	N	Sterilization of health care products_ - Ethylene oxide_ - Part_1: Requirements for development, validation and routine control of a sterilization process for medical devices	Y				

ISO 11137-1	2006-04	N	Sterilization of health care products_- Radiation_- Part_1: Requirements for development, validation and routine control of a sterilization process for medical devices	Y				
ISO 11137-2	2012-03	N	Sterilization of health care products_- Radiation_- Part_2: Establishing the sterilization dose	Y				
ISO 11137-3	2006-04	N	Sterilization of health care products_- Radiation_- Part_3: Guidance on dosimetric aspects	N				
ISO 11138-1	2006-07	N	Sterilization of health care products_- Biological indicators_- Part_1: General requirements	N				
ISO 11138-2	2006-07	N	Sterilization of health care products_- Biological indicators_- Part_2: Biological indicators for ethylene oxide sterilization processes	Y				
ISO 11138-3	2006-07	N	Sterilization of health care products_- Biological indicators_- Part_3: Biological indicators for moist heat sterilization processes	Y				
ISO 11138-4	2006-07	N	Sterilization of health care products_- Biological indicators_- Part_4: Biological indicators for dry heat sterilization processes	N				
ISO 11138-5	2006-07	N	Sterilization of health care products_- Biological indicators_- Part_5: Biological indicators for low-temperature steam and formaldehyde sterilization processes	N				
ISO 11140-1	2005-07	N	Sterilization of health care products_- Chemical indicators_- Part_1: General requirements	Y				
ISO 11140-3	2007-03	N	Sterilization of health care products_- Chemical indicators_- Part_3: Class_2 indicator systems for use in the Bowie and Dick-type steam penetration test	Y				
ISO 11140-3 Technical	2007-11	N	Sterilization of health care products_- Chemical indicators_- Part_3: Class 2 indicator systems for use in the Bowie and Dick-type steam penetration test; Technical Corrigendum_1	Y				
ISO 11140-4	2007-03	N	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 penetration	N				

ISO 11140-5	2007-03	N	Sterilization of health care products_ - Chemical indicators_ - Part_5: Class_2 indicators for Bowie and Dick-type air removal tests	N				
ISO 11143	2008-07	N	Dentistry_ - Amalgam separators					
ISO 11144	1995-05	N	Dental equipment_ - Connections for supply and waste lines					
ISO 11156	2011-07	N	Packaging_ - Accessible design_ - General requirements					
ISO 11193-1	2008-09	N	Single-use medical examination gloves_ - Part_1: Specification for gloves made from rubber latex or rubber solution	N				
ISO 11193-2	2006-11	N	Single-use medical examination gloves_ - Part_2: Specification for gloves made from poly(vinyl chloride)	N				
ISO 11195	1995-10	N	Gas mixers for medical use_ - Stand-alone gas mixers	N				
ISO 11197	2004-12	N	Medical supply units	Y				
ISO 11199-1	1999-08	N	Walking aids manipulated by both arms_ - Requirements and test methods_ - Part_1: Walking frames					
ISO 11199-2	2005-04	N	Walking aids manipulated by both arms_ - Requirements and test methods_ - Part_2: Rollators					
ISO 11199-3	2005-04	N	Walking aids manipulated by both arms_ - Requirements and test methods_ - Part_3: Walking tables					
ISO 11318	2002-08	N	Cardiac defibrillators_ - Connector assembly DF-1 for implantable defibrillators_ - Dimensional and test requirements	N				
ISO 11334-1	2007-02	N	Assistive products for walking manipulated by one arm_ - Requirements and test methods_ - Part_1: Elbow crutches					
ISO 11334-4	1999-02	N	Walking aids manipulated by one arm_ - Requirements and test methods_ - Part_4: Walking sticks with three or more legs					
ISO 1135-3	1986-11	N	Transfusion equipment for medical use; Part 3 : Blood-taking set	N				
ISO 1135-4	2012-03	N	Transfusion equipment for medical use_ - Part_4: Transfusion sets for single use	Y				
ISO 11380	1994-10	N	Optics and optical instruments_ - Ophthalmic optics_ - Formers	N				
ISO 11381	1994-12	N	Optics and optical instruments_ - Ophthalmic optics_ - Screw threads	N				
ISO 11418-1	2005-02	N	Containers and accessories for pharmaceutical preparations_ - Part_1: Drop-dispensing glass bottles					

ISO 11418-2	2005-02	N	Containers and accessories for pharmaceutical preparations_- Part_2: Screw-neck glass bottles for syrups					
ISO 11418-3	2005-02	N	Containers and accessories for pharmaceutical preparations_- Part_3: Screw-neck glass bottles (veral) for solid and liquid dosage forms					
ISO 11418-4	2005-02	N	Containers and accessories for pharmaceutical preparations_- Part_4: Tablet glass bottles					
ISO 11418-5	1997-12	N	Containers and accessories for pharmaceutical preparations_- Part_5: Dropper assemblies					
ISO 11418-7	1998-10	N	Containers and accessories for pharmaceutical preparations_- Part_7: Screw-neck vials made of glass tubing for liquid dosage forms					
ISO 11498	1997-02	N	Dental handpieces_- Dental low-voltage electrical motors					
ISO 11499	2007-07	N	Dentistry_- Single-use cartridges for local anaesthetics	N				
ISO 11607-1	2006-04	N	Packaging for terminally sterilized medical devices_- Part_1: Requirements for materials, sterile barrier systems and packaging systems	Y				
ISO 11607-2	2006-04	N	Packaging for terminally sterilized medical devices_- Part_2: Validation requirements for forming, sealing and assembly processes	Y				
ISO 11608-1	2000-12	N	Pen-injectors for medical use_- Part_1: Pen-injectors; Requirements and test methods	N				
ISO 11608-2	2000-12	N	Pen-injectors for medical use_- Part_2: Needles; Requirements and test methods	N				
ISO 11608-3	2000-12	N	Pen-injectors for medical use_- Part_3: Finished cartridges; Requirements and test methods	N				
ISO 11608-4	2006-03	N	Pen-injectors for medical use_- Part_4: Requirements and test methods for electronic and electromechanical pen-injectors	N				
ISO 11609	2010-09	N	Dentistry_- Dentifrices_- Requirements, test methods and marking					
ISO 11663	2009-04	N	Quality of dialysis fluid for haemodialysis and related therapies	N				
ISO 11683	1997-10	N	Packaging_- Tactile warnings of danger_- Requirements					

ISO 11712	2009-05	N	Anaesthetic and respiratory equipment_- Supralaryngeal airways and connectors	N				
ISO 11737-1	2006-04	N	Sterilization of medical devices_- Microbiological methods_- Part_1: Determination of a population of microorganisms on products	Y				
ISO 11737-1 Technical	2007-05	N	Sterilization of medical devices_- Microbiological methods_- Part_1: Determination of a population of microorganisms on products; Technical Corrigendum_1	Y				
ISO 11737-2	2009-11	N	Sterilization of medical devices_- Microbiological methods_- Part_2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process	Y				
ISO 11810-1	2005-02	N	Lasers and laser-related equipment_- Test method and classification for the laser resistance of surgical drapes and/or patient protective covers_- Part_1: Primary ignition and penetration	Y				
ISO 11810-2	2007-05	N	Lasers and laser-related equipment_- Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers_- Part_2: Secondary ignition	Y				
ISO 11904-1	2002-10	N	Acoustics_- Determination of sound immission from sound sources placed close for the ear_- Part_1: Technique using a microphone in a real ear (MIRE technique)					
ISO 11948-1	1996-11	N	Urine-absorbing aids_- Part_1: Whole-product testing					
ISO 11953	2010-06	N	Dentistry_- Implants_- Clinical performance of hand torque instruments					
ISO 11978	2000-03	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Information supplied by the manufacturer	N				
ISO 11979-1	2006-07	N	Ophthalmic implants_- Intraocular lenses_- Part_1: Vocabulary	N				
ISO 11979-10	2006-08	N	Ophthalmic implants_- Intraocular lenses_- Part_10: Phakic intraocular lenses	N				
ISO 11979-2	1999-12	N	Ophthalmic implants_- Intraocular lenses_- Part_2: Optical properties and test methods	N				

ISO 11979-2 Technical	2003-11	N	Ophthalmic implants_- Intraocular lenses_- Part_2: Optical properties and test methods; Technical Corrigendum_1	N				
ISO 11979-3	2006-05	N	Ophthalmic implants_- Intraocular lenses_- Part_3: Mechanical properties and test methods	N				
ISO 11979-4	2008-12	N	Ophthalmic implants_- Intraocular lenses_- Part_4: Labelling and information	N				
ISO 11979-5	2006-06	N	Ophthalmic implants_- Intraocular lenses_- Part_5: Biocompatibility	N				
ISO 11979-6	2007-07	N	Ophthalmic implants_- Intraocular lenses_- Part_6: Shelf-life and transport stability	N				
ISO 11979-7	2006-05	N	Ophthalmic implants_- Intraocular lenses_- Part_7: Clinical investigations	N				
ISO 11979-7 AMD 1	2012-01	N	Ophthalmic implants_- Intraocular lenses_- Part_7: Clinical investigations; Amendment_1	N				
ISO 11979-8	2006-07	N	Ophthalmic implants_- Intraocular lenses_- Part_8: Fundamental requirements	Y				
ISO 11979-8 AMD 1	2011-05	N	Ophthalmic implants_- Intraocular lenses_- Part_8: Fundamental requirements; Amendment_1	Y				
ISO 11979-9	2006-09	N	Ophthalmic implants_- Intraocular lenses_- Part_9: Multifocal intraocular lenses	N				
ISO 11980	2009-10	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Guidance for clinical investigations	N				
ISO 11981	2009-07	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Determination of physical compatibility of contact lens care products with contact lenses	N				
ISO 11985	1997-12	N	Ophthalmic optics_- Contact lenses_- Ageing by exposure to UV and visible radiation (in vitro method)	N				
ISO 11986	2010-11	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Determination of preservative uptake and release	N				
ISO 11987	1997-12	N	Ophthalmic optics_- Contact lenses_- Determination of shelf-life	N				

ISO 11987 Technical C	1998-04	N	Ophthalmic optics_- Contact lenses_- Determination of shelf-life; Technical Corrigendum_1	N				
ISO 11990-1	2011-08	N	Lasers and laser-related equipment_- Determination of laser resistance of tracheal tubes_- Part_1: Tracheal tube shaft	Y if proposed by CEN				
ISO 11990-2	2010-07	N	Lasers and laser-related equipment_- Determination of laser resistance of tracheal tubes_- Part_2: Tracheal tube cuffs	Y if proposed by CEN				
ISO 12052	2006-11	N	Health informatics_- Digital imaging and communication in medicine (DICOM) including workflow and data management					
ISO 12124	2001-03	N	Acoustics_- Procedures for the measurement of real-ear acoustical characteristics of hearing aids					
ISO 12189	2008-05	N	Implants for surgery_- Mechanical testing of implantable spinal devices_- Fatigue test method for spinal implant assemblies using an anterior support	N				
ISO 12243	2003-10	N	Medical gloves made from natural rubber latex_- Determination of water-extractable protein using the modified Lowry method	N				
ISO 12625-1	2011-08	N	Tissue paper and tissue products_- Part_1: General guidance on terms					
ISO 12625-12	2010-01	N	Tissue paper and tissue products_- Part_12: Determination of tensile strength of perforated lines_- Calculation of perforation efficiency					
ISO 12625-3	2005-04	N	Tissue paper and tissue products_- Part_3: Determination of thickness, bulking thickness and apparent bulk density					
ISO 12625-4	2005-04	N	Tissue paper and tissue products_- Part_4: Determination of tensile strength, stretch at break and tensile energy absorption					
ISO 12625-5	2005-04	N	Tissue paper and tissue products_- Part_5: Determination of wet tensile strength					
ISO 12625-6	2005-02	N	Tissue paper and tissue products_- Part_6: Determination of grammage					
ISO 12625-7	2007-03	N	Tissue paper and tissue products_- Part_7: Determination of optical properties					
ISO 12625-8	2010-12	N	Tissue paper and tissue products_- Part_8: Water-absorption time and water-absorption capacity, basket-immersion test method					
ISO 12625-9	2005-05	N	Tissue paper and tissue products_- Part_9: Determination of ball burst strength					
ISO 12864	1997-12	N	Ophthalmic optics_- Contact lenses_- Determination of scattered light	N				

ISO 12865	2006-07	N	Ophthalmic instruments - Retinoscopes	N				
ISO 12866	1999-06	N	Ophthalmic instruments - Perimeters	N				
ISO 12866 AMD 1	2008-11	N	Ophthalmic instruments - Perimeters; Amendment_1	N				
ISO 12867	2010-06	N	Ophthalmic instruments - Trial frames	N				
ISO 12870	2004-08	N	Ophthalmic optics - Spectacle frames - Requirements and test methods	Y				
ISO 12891-1	2011-05	N	Implants for surgery - Retrieval and analysis of surgical implants - Part_1: Retrieval and handling	N				
ISO 12891-2	2000-02	N	Retrieval and analysis of surgical implants - Part_2: Analysis of retrieved metallic surgical implants	N				
ISO 12891-3	2000-02	N	Retrieval and analysis of surgical implants - Part_3: Analysis of retrieved polymeric surgical implants	N				
ISO 12891-4	2000-02	N	Retrieval and analysis of surgical implants - Part_4: Analysis of retrieved ceramic surgical implants	N				
ISO 12967-1	2009-08	N	Health informatics - Service architecture - Part_1: Enterprise viewpoint					
ISO 12967-2	2009-08	N	Health informatics - Service architecture - Part_2: Information viewpoint					
ISO 12967-3	2009-08	N	Health informatics - Service architecture - Part_3: Computational viewpoint					
ISO 13212	2011-05	N	Ophthalmic optics - Contact lens care products - Guidelines for determination of shelf-life	N				
ISO 13294	1997-05	N	Dental handpieces - Dental air-motors					
ISO 13295	2007-07	N	Dentistry - Mandrels for rotary instruments					
ISO 13356	2008-06	N	Implants for surgery - Ceramic materials based on yttria-stabilized tetragonal zirconia (Y-TZP)	N				
ISO 13397-1	1995-12	N	Periodontal curettes, dental scalers and excavators - Part_1: General requirements					
ISO 13397-2	2005-06	N	Dentistry - Periodontal curettes, dental scalers and excavators - Part_2: Periodontal curettes of Gr-type					
ISO 13397-3	1996-09	N	Periodontal curettes, dental scalers and excavators - Part_3: Dental scalers - H-type					
ISO 13397-4	1997-12	N	Periodontal curettes, dental scalers and excavators - Part_4: Dental excavators - Discoid type					
ISO 13402	1995-08	N	Surgical and dental hand instruments - Determination of resistance against autoclaving, corrosion and thermal exposure	N				

ISO 13404	2007-07	N	Prosthetics and orthotics_ - Categorization and description of external orthoses and orthotic components					
ISO 13405-1	1996-10	N	Prosthetics and orthotics_ - Classification and description of prosthetic components_ - Part_1: Classification of prosthetic components					
ISO 13405-2	1996-10	N	Prosthetics and orthotics_ - Classification and description of prosthetic components_ - Part_2: Description of lower-limb prosthetic components					
ISO 13405-3	1996-10	N	Prosthetics and orthotics_ - Classification and description of prosthetic components_ - Part_3: Description of upper-limb prosthetic components					
ISO 13408-1	2008-06	N	Aseptic processing of health care products_ - Part_1: General requirements	Y				
ISO 13408-2	2003-03	N	Aseptic processing of health care products_ - Part_2: Filtration	Y				
ISO 13408-3	2006-09	N	Aseptic processing of health care products_ - Part_3: Lyophilization	Y				
ISO 13408-4	2005-11	N	Aseptic processing of health care products_ - Part_4: Clean-in-place technologies	Y				
ISO 13408-5	2006-11	N	Aseptic processing of health care products_ - Part_5: Sterilization in place	Y				
ISO 13408-6	2005-06	N	Aseptic processing of health care products_ - Part_6: Isolator systems	Y				
ISO 13485	2003-07	N	Medical devices_ - Quality management systems_ - Requirements for regulatory purposes	P				
ISO 13485 Technical C	2009-08	N	Medical devices_ - Quality management systems_ - Requirements for regulatory purposes; Technical Corrigendum_1	P				
ISO 13606-1	2008-02	N	Health informatics_ - Electronic health record communication_ - Part_1: Reference model					
ISO 13606-2	2008-12	N	Health informatics_ - Electronic health record communication_ - Part_2: Archetype interchange specification					
ISO 13606-3	2009-02	N	Health informatics_ - Electronic health record communication_ - Part_3: Reference archetypes and term lists					
ISO 13606-5	2010-03	N	Health informatics_ - Electronic health record communication_ - Part_5: Interface specification					
ISO 13666	1998-08	N	Ophthalmic optics_ - Spectacle lenses_ - Vocabulary	N				

ISO 13716	1999-05	N	Dentistry_- Reversible-irreversible hydrocolloid impression material systems					
ISO 13779-1	2008-10	N	Implants for surgery_- Hydroxyapatite_- Part_1: Ceramic hydroxyapatite	N				
ISO 13779-2	2008-10	N	Implants for surgery_- Hydroxyapatite_- Part_2: Coatings of hydroxyapatite	N				
ISO 13779-3	2008-02	N	Implants for surgery_- Hydroxyapatite_- Part_3: Chemical analysis and characterization of crystallinity and phase purity	N				
ISO 13779-4	2002-05	N	Implants for surgery_- Hydroxyapatite_- Part_4: Determination of coating adhesion strength	N				
ISO 13781	1997-02	N	Poly(L-lactide) resins and fabricated forms for surgical implants_- In vitro degradation testing	N				
ISO 13782	1996-12	N	Implants for surgery_- Metallic materials_- Unalloyed tantalum for surgical implant applications	N				
ISO 13897	2003-02	N	Dentistry_- Amalgam capsules					
ISO 13897 Technical Co	2003-12	N	Dentistry_- Amalgam capsules; Technical Corrigendum_1					
ISO 13926-1	2004-11	N	Pen systems_- Part_1: Glass cylinders for pen-injectors for medical use	N				
ISO 13926-2	2011-04	N	Pen systems_- Part_2: Plunger stoppers for pen-injectors for medical use	N				
ISO 13958	2009-04	N	Concentrates for haemodialysis and related therapies	N				
ISO 13959	2009-04	N	Water for haemodialysis and related therapies	N				
ISO 13960	2010-07	N	Cardiovascular implants and extracorporeal systems_- Plasmafilters	N				
ISO 14155	2011-02	N	Clinical investigation of medical devices for human subjects_- Good clinical practice	Y				
ISO 14155 Technical C	2011-07	N	Clinical investigation of medical devices for human subjects_- Good clinical practice; Technical Corrigendum_1	Y				

ISO 14160	2011-07	N	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	Y				
ISO 14161	2009-09	N	Sterilization of health care products_ - Biological indicators_ - Guidance for the selection, use and interpretation of results	N				
ISO 14233	2003-03	N	Dentistry_ - Polymer-based die materials					
ISO 14242-1	2012-01	N	Implants for surgery_ - Wear of total hip-joint prostheses_ - Part_1: Loading and displacement parameters for wear-testing machines and corresponding environmental conditions for test	N				
ISO 14242-2	2000-09	N	Implants for surgery_ - Wear of total hip joint prostheses_ - Part_2: Methods of measurement	N				
ISO 14242-3	2009-03	N	Implants for surgery_ - Wear of total hip-joint prostheses_ - Part_3: Loading and displacement parameters for orbital bearing type wear testing machines and corresponding environmental conditions for test	N				
ISO 14243-1	2009-11	N	Implants for surgery_ - Wear of total knee-joint prostheses_ - Part_1: Loading and displacement parameters for wear-testing machines with load control and corresponding environmental conditions for test	N				
ISO 14243-2	2009-11	N	Implants for surgery_ - Wear of total knee-joint prostheses_ - Part_2: Methods of measurement	N				
ISO 14243-3	2004-09	N	Implants for surgery_ - Wear of total knee-joint prostheses_ - Part_3: Loading and displacement parameters for wear-testing machines with displacement control and corresponding environmental conditions for test	N				

ISO 14243-3 Technical	2006-02	N	Implants for surgery_- Wear of total knee-joint prostheses_- Part_3: Loading and displacement parameters for wear-testing machines with displacement control and corresponding environmental conditions for test	N				
ISO 14356	2003-03	N	Dentistry_- Duplicating material					
ISO 14408	2005-06	N	Tracheal tubes designed for laser surgery_- Requirements for marking and accompanying information	Y				
ISO 14534	2011-04	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Fundamental requirements	Y				
ISO 14602	2010-04	N	Non-active surgical implants_- Implants for osteosynthesis_- Particular requirements	Y				
ISO 14607	2007-02	N	Non-active surgical implants_- Mammary implants_- Particular requirements	Y				
ISO 14630	2008-01	N	Non-active surgical implants_- General requirements	Y				
ISO 14708-1	2000-11	N	Implants for surgery_- Active implantable medical devices_- Part_1: General requirements for safety, marking and for information to be provided by the manufacturer	N				
ISO 14708-2	2005-10	N	Implants for surgery_- Active implantable medical devices_- Part_2: Cardiac pacemakers	N				
ISO 14708-3	2008-11	N	Implants for surgery_- Active implantable medical devices_- Part_3: Implantable neurostimulators	N				
ISO 14708-4	2008-11	N	Implants for surgery_- Active implantable medical devices_- Part_4: Implantable infusion pumps	N				
ISO 14708-5	2010-02	N	Implants for surgery_- Active implantable medical devices_- Part_5: Circulatory support devices	N				
ISO 14708-6	2010-03	N	Implants for surgery_- Active implantable medical devices_- Part_6: Particular requirements for active implantable medical devices intended to treat tachyarrhythmia (including implantable defibrillators)	N				

ISO 14729	2001-04	N	Ophthalmic optics_ - Contact lens care products_ - Microbiological requirements and test methods for products and regimens for hygienic management of contact lenses	N				
ISO 14729 AMD 1	2010-10	N	Ophthalmic optics_ - Contact lens care products_ - Microbiological requirements and test methods for products and regimens for hygienic management of contact lenses; Amendment_1	N				
ISO 14730	2000-09	N	Ophthalmic optics_ - Contact lens care products_ - Antimicrobial preservative efficacy testing and guidance on determining discard date	N				
ISO 14801	2007-11	N	Dentistry_ - Implants_ - Dynamic fatigue test for endosseous dental implants					
ISO 14879-1	2000-06	N	Implants for surgery_ - Total knee-joint prostheses_ - Part_1: Determination of endurance properties of knee tibial trays	N				
ISO 14889	2003-05	N	Ophthalmic optics_ - Spectacle lenses_ - Fundamental requirements for uncut finished lenses	Y				
ISO 14937	2009-10	N	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	Y				
ISO 14949	2001-10	N	Implants for surgery_ - Two-part addition-cure silicone elastomers	N				
ISO 14971	2007-03	N	Medical devices_ - Application of risk management to medical devices	P				
ISO 14972	1998-12	N	Sterile obturators for single use with over-needle peripheral intravascular catheters	N				
ISO 15001	2010-06	N	Anaesthetic and respiratory equipment_ - Compatibility with oxygen	Y				
ISO 15002	2008-07	N	Flow-metering devices for connection to terminal units of medical gas pipeline systems	Y				
ISO 15004-1	2006-06	N	Ophthalmic instruments_ - Fundamental requirements and test methods_ - Part_1: General requirements applicable to all ophthalmic instruments	Y				

ISO 15004-2	2007-02	N	Ophthalmic instruments_ - Fundamental requirements and test methods_ - Part_2: Light hazard protection	N				
ISO 15010	1998-06	N	Disposable hanging devices for transfusion and infusion bottles_ - Requirements and test methods	N				
ISO 15032	2000-04	N	Prostheses_ - Structural testing of hip units	N				
ISO 15087-1	1999-11	N	Dental elevators_ - Part_1: General requirements					
ISO 15087-2	2000-04	N	Dental elevators_ - Part_2: Warwick James elevators					
ISO 15087-3	2000-05	N	Dental elevators_ - Part_3: Cryer elevators					
ISO 15087-4	2000-05	N	Dental elevators_ - Part_4: Coupland elevators					
ISO 15087-5	2000-05	N	Dental elevators_ - Part_5: Bein elevators					
ISO 15087-6	2000-05	N	Dental elevators_ - Part_6: Flohr elevators					
ISO 15098-1	1999-10	N	Dental tweezers_ - Part_1: General requirements					
ISO 15098-2	2000-02	N	Dental tweezers_ - Part_2: Meriam types					
ISO 15098-3	2000-02	N	Dental tweezers_ - Part_3: College types					
ISO 15137	2005-07	N	Self-adhesive hanging devices for infusion bottles and injection vials_ - Requirements and test methods	N				
ISO 15142-1	2003-08	N	Implants for surgery_ - Metal intramedullary nailing systems_ - Part_1: Intramedullary nails	N				
ISO 15142-2	2003-08	N	Implants for surgery_ - Metal intramedullary nailing systems_ - Part_2: Locking components	N				
ISO 15142-3	2003-08	N	Implants for surgery_ - Metal intramedullary nailing systems_ - Part_3: Connection devices and reamer diameter measurements	N				
ISO 15193	2009-05	N	In vitro diagnostic medical devices_ - Measurement of quantities in samples of biological origin_ - Requirements for content and presentation of reference measurement procedures	Y				
ISO 15194	2009-05	N	In vitro diagnostic medical devices_ - Measurement of quantities in samples of biological origin_ - Requirements for certified reference materials and the content of supporting documentation	Y				
ISO 15197	2003-05	N	In vitro diagnostic test systems_ - Requirements for blood-glucose monitoring systems for self-testing in managing diabetes mellitus	N				

ISO 15198	2004-07	N	Clinical laboratory medicine_- In vitro diagnostic medical devices_- Validation of user quality control procedures by the manufacturer	N				
ISO 15223-1	2007-04	N	Medical devices_- Symbols to be used with medical device labels, labelling and information to be supplied_- Part_1: General requirements	N				
ISO 15223-1 AMD 1	2008-06	N	Medical devices_- Symbols to be used with medical device labels, labelling and information to be supplied_- Part_1: General requirements; Amendment_1	N				
ISO 15223-2	2010-01	N	Medical devices_- Symbols to be used with medical device labels, labelling, and information to be supplied_- Part_2: Symbol development, selection and validation	N				
ISO 15225	2010-05	N	Medical devices_- Quality management_- Medical device nomenclature data structure	N				
ISO 15253	2000-09	N	Ophthalmic optics and instruments_- Optical devices for enhancing low vision					
ISO 15254	2009-07	N	Ophthalmic optics and instruments_- Electro-optical devices for enhancing low vision	N				
ISO 15374	1998-08	N	Implants for surgery_- Requirements for production of forgings	N				
ISO 15375	2010-06	N	Medical infusion bottles_- Suspension devices for multiple use_- Requirements and test methods	N				
ISO 15378	2011-11	N	Primary packaging materials for medicinal products_- Particular requirements for the application of ISO_9001:2008, with reference to Good Manufacturing Practice_(GMP)					
ISO 15606	1999-12	N	Dental handpieces_- Air-powered scalers and scaler tips					
ISO 15621	2011-02	N	Urine-absorbing aids_- General guidelines on evaluation					
ISO 1563	1990-09	N	Dental alginate impression material					
ISO 1564	1995-11	N	Dental aqueous impression materials based on agar					
ISO 15674	2009-04	N	Cardiovascular implants and artificial organs_- Hard-shell cardiotomy/venous reservoir systems (with/without filter) and soft venous reservoir bags	N				

ISO 15675	2009-04	N	Cardiovascular implants and artificial organs_- Cardiopulmonary bypass systems_- Arterial blood line filters	N				
ISO 15676	2005-07	N	Cardiovascular implants and artificial organs_- Requirements for single-use tubing packs for cardiopulmonary bypass and extracorporeal membrane oxygenation (ECMO)	N				
ISO 15747	2010-04	N	Plastic containers for intravenous injections	Y				
ISO 15752	2010-01	N	Ophthalmic instruments_- Endoilluminators_- Fundamental requirements and test methods for optical radiation safety	N				
ISO 15759	2005-04	N	Medical infusion equipment_- Plastics caps with inserted elastomeric liner for containers manufactured by the blow- fill-seal (BFS) process	N				
ISO 15798	2010-01	N	Ophthalmic implants_- Ophthalmic viscosurgical devices	Y				
ISO 15814	1999-11	N	Implants for surgery_- Copolymers and blends based on polylactide - In vitro degradation testing	N				
ISO 15841	2006-10	N	Dentistry_- Wires for use in orthodontics					
ISO 15854	2005-07	N	Dentistry_- Casting and baseplate waxes					
ISO 15882	2008-09	N	Sterilization of health care products_- Chemical indicators_- Guidance for selection, use and interpretation of results	N				
ISO 15883-1	2006-04	N	Washer-disinfectors_- Part_1: General requirements, terms and definitions and tests	Y				
ISO 15883-2	2006-04	N	Washer-disinfectors_- Part_2: Requirements and tests for washer-disinfectors employing thermal disinfection for surgical instruments, anaesthetic equipment, bowls, dishes, receivers, utensils, glassware, etc.	Y				
ISO 15883-3	2006-04	N	Washer-disinfectors_- Part_3: Requirements and tests for washer-disinfectors employing thermal disinfection for human waste containers	Y				
ISO 15883-4	2008-05	N	Washer-disinfectors_- Part_4: Requirements and tests for washer-disinfectors employing chemical disinfection for thermolabile endoscopes	Y				

ISO 15883-6	2011-04	N	Washer-disinfectors_- Part_6: Requirements and tests for washer-disinfectors employing thermal disinfection for non-invasive, non-critical medical devices and healthcare equipment	Y if proposed by CEN				
ISO 15912	2006-10	N	Dentistry_- Casting investments and refractory die materials					
ISO 15912 AMD 1	2011-07	N	Dentistry_- Casting investments and refractory die materials; Amendment_1: Requirement and test method for adequacy of expansion of Type_1 and Type_2 materials					
ISO 16021	2000-11	N	Urine-absorbing aids_- Basic principles for evaluation of single-use adult-incontinence-absorbing aids from the perspective of users and caregivers					
ISO 16034	2002-02	N	Ophthalmic optics_- Specifications for single-vision ready-to-wear near-vision spectacles	N				
ISO 16034 Technical C	2006-08	N	Ophthalmic optics_- Specifications for single-vision ready-to-wear near- vision spectacles; Technical Corrigendum_1	N				
ISO 16037	2002-05	N	Rubber condoms for clinical trials_- Measurement of physical properties					
ISO 16037 AMD 1	2011-02	N	Rubber condoms for clinical trials_- Measurement of physical properties; Amendment_1					
ISO 16038	2005-11	N	Rubber condoms_- Guidance on the use of ISO_4074 in the quality management of natural rubber latex condoms					
ISO 16054	2000-12	N	Implants for surgery_- Minimum data sets for surgical implants	N				
ISO 16059	2007-08	N	Dentistry_- Required elements for codification used in data exchange					
ISO 16061	2008-12	N	Instrumentation for use in association with non-active surgical implants_- General requirements	Y				
ISO 16201	2006-10	N	Technical aids for persons with disability_- Environmental control systems for daily living	Y				
ISO 16284	2006-03	N	Ophthalmic optics_- Information interchange for ophthalmic optical equipment	N				
ISO 16391	2002-10	N	Aids for ostomy and incontinence_- Irrigation sets_- Requirements and test methods					

ISO 16402	2008-05	N	Implants for surgery_ - Acrylic resin cement_ - Flexural fatigue testing of acrylic resin cements used in orthopaedics	N				
ISO 16408	2004-04	N	Dentistry_ - Oral hygiene products_ - Oral rinses					
ISO 16409	2006-10	N	Dentistry_ - Oral hygiene products_ - Manual interdental brushes					
ISO 16409 AMD 1	2010-02	N	Dentistry_ - Oral hygiene products_ - Manual interdental brushes; Amendment_1					
ISO 16428	2005-04	N	Implants for surgery_ - Test solutions and environmental conditions for static and dynamic corrosion tests on implantable materials and medical devices	N				
ISO 16429	2004-07	N	Implants for surgery_ - Measurements of open-circuit potential to assess corrosion behaviour of metallic implantable materials and medical devices over extended time periods	N				
ISO 16628	2008-11	N	Tracheobronchial tubes_ - Sizing and marking	N				
ISO 16671	2003-05	N	Ophthalmic implants_- Irrigating solutions for ophthalmic surgery	N				
ISO 16672	2003-02	N	Ophthalmic implants_- Ocular endotamponades	N				
ISO 16840-1	2006-03	N	Wheelchair seating_ - Part_1: Vocabulary, reference axis convention and measures for body segments, posture and postural support surfaces					
ISO 16840-2	2007-07	N	Wheelchair seating_ - Part_2: Determination of physical and mechanical characteristics of devices intended to manage tissue integrity_- Seat cushions					
ISO 16840-3	2006-07	N	Wheelchair seating_ - Part_3: Determination of static, impact and repetitive load strengths for postural support devices					
ISO 16840-4	2009-03	N	Wheelchair seating_ - Part_4: Seating systems for use in motor vehicles					
ISO 17090-1	2008-02	N	Health informatics_ - Public key infrastructure_- Part_1: Overview of digital certificate services					
ISO 17090-2	2008-02	N	Health informatics_ - Public key infrastructure_- Part_2: Certificate profile					
ISO 17090-3	2008-02	N	Health informatics_ - Public key infrastructure_- Part_3: Policy management of certification authority					
ISO 17115	2007-07	N	Health informatics_ - Vocabulary for terminological systems					

ISO 17190-1	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_1: Determination of pH					
ISO 17190-10	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_10: Determination of extractable polymer content by potentiometric titration					
ISO 17190-11	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_11: Determination of content of respirable particles					
ISO 17190-2	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_2: Determination of amount of residual monomers					
ISO 17190-3	2001-12	N	Urine absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_3: Determination of particle size distribution by sieve fractionation					
ISO 17190-4	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_4: Determination of moisture content by mass loss upon heating					
ISO 17190-5	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_5: Gravimetric determination of free swell capacity in saline solution					
ISO 17190-6	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_6: Gravimetric determination of fluid retention capacity in saline solution after centrifugation					
ISO 17190-7	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_7: Gravimetric determination of absorption under pressure					
ISO 17190-8	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_8: Gravimetric determination of flowrate					
ISO 17190-9	2001-12	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_9: Gravimetric determination of density					

ISO 17190-9 Technical C	2002-10	N	Urine-absorbing aids for incontinence_- Test methods for characterizing polymer-based absorbent materials_- Part_9: Gravimetric determination of density; Technical Corrigendum_1					
ISO 17191	2004-02	N	Urine-absorbing aids for incontinence_- Measurement of airborne respirable polyacrylate superabsorbent materials_- Determination of dust in collection cassettes by sodium atomic absorption spectrometry					
ISO 17432	2004-12	N	Health informatics_- Messages and communication_- Web access to DICOM persistent objects					
ISO 17510-1	2007-10	N	Sleep apnoea breathing therapy_- Part_1: Sleep apnoea breathing therapy equipment	Y				
ISO 17510-2	2007-10	N	Sleep apnoea breathing therapy_- Part_2: Masks and application accessories	Y				
ISO 17511	2003-08	N	In vitro diagnostic medical devices_- Measurement of quantities in biological samples_- Metrological traceability of values assigned to calibrators and control materials	Y				
ISO 17593	2007-04	N	Clinical laboratory testing and in vitro medical devices_- Requirements for in vitro monitoring systems for self-testing of oral anticoagulant therapy	N				
ISO 17664	2004-03	N	Sterilization of medical devices_- Information to be provided by the manufacturer for the processing of resterilizable medical devices	Y				
ISO 17665-1	2006-08	N	Sterilization of health care products_- Moist heat_- Part_1: Requirements for the development, validation and routine control of a sterilization process for medical devices	Y				
ISO 17853	2011-03	N	Wear of implant materials_- Polymer and metal wear particles_- Isolation and characterization	N				
ISO 1797-1	2011-08	N	Dentistry_- Shanks for rotary instruments_- Part_1: Shanks made of metals					
ISO 1797-2	1992-02	N	Dental rotary instruments; shanks; part_2: shanks made of plastics					
ISO 18084	2011-09	N	Press tools for tablets_- Punches and dies					
ISO 18104	2003-12	N	Health informatics_- Integration of a reference terminology model for nursing					

ISO 18113-1	2009-12	N	In vitro diagnostic medical devices_ - Information supplied by the manufacturer (labelling)_ - Part_1: Terms, definitions and general requirements	Y				
ISO 18113-2	2009-12	N	In vitro diagnostic medical devices_ - Information supplied by the manufacturer (labelling)_ - Part_2: In vitro diagnostic reagents for professional use	Y				
ISO 18113-3	2009-12	N	In vitro diagnostic medical devices_ - Information supplied by the manufacturer (labelling)_ - Part_3: In vitro diagnostic instruments for professional use	Y				
ISO 18113-4	2009-12	N	In vitro diagnostic medical devices_ - Information supplied by the manufacturer (labelling)_ - Part_4: In vitro diagnostic reagents for self-testing	Y				
ISO 18113-5	2009-12	N	In vitro diagnostic medical devices_ - Information supplied by the manufacturer (labelling)_ - Part_5: In vitro diagnostic instruments for self-testing	Y				
ISO 18153	2003-08	N	In vitro diagnostic medical devices_ - Measurement of quantities in biological samples_ - Metrological traceability of values for catalytic concentration of enzymes assigned calibrators and control materials	Y				
ISO 18192-1	2011-03	N	Implants for surgery_ - Wear of total intervertebral spinal disc prostheses_ - Part_1: Loading and displacement parameters for wear testing and corresponding environmental conditions for test	N				
ISO 18192-2	2010-06	N	Implants for surgery_ - Wear of total intervertebral spinal disc prostheses_ - Part_2: Nucleus replacements	N				
ISO 18232	2006-04	N	Health Informatics_ - Messages and communication_ - Format of length limited globally unique string identifiers					
ISO 18308	2011-04	N	Health informatics_ - Requirements for an electronic health record architecture					
ISO 18369-1	2006-08	N	Ophthalmic optics_ - Contact lenses_ - Part_1: Vocabulary, classification system and recommendations for labelling specifications	N				

ISO 18369-1 AMD 1	2009-02	N	Ophthalmic optics - Contact lenses - Part 1: Vocabulary, classification system and recommendations for labelling specifications; Amendment 1	N				
ISO 18369-2	2006-08	N	Ophthalmic optics - Contact lenses - Part 2: Tolerances	N				
ISO 18369-3	2006-08	N	Ophthalmic optics - Contact lenses - Part 3: Measurement methods	N				
ISO 18369-4	2006-08	N	Ophthalmic optics - Contact lenses - Part 4: Physicochemical properties of contact lens materials	N				
ISO 18472	2006-06	N	Sterilization of health care products - Biological and chemical indicators - Test equipment	N				
ISO 18777	2005-02	N	Transportable liquid oxygen systems for medical use - Particular requirements	Y				
ISO 18778	2005-02	N	Respiratory equipment - Infant monitors - Particular requirements	Y				
ISO 18779	2005-02	N	Medical devices for conserving oxygen and oxygen mixtures - Particular requirements	Y				
ISO 18812	2003-03	N	Health informatics - Clinical analyser interfaces to laboratory information systems - Use profiles					
ISO 19001	2002-11	N	In vitro diagnostic medical devices - Information supplied by the manufacturer with in vitro diagnostic reagents for staining in biology	N				
ISO 19054	2005-07	N	Rail systems for supporting medical equipment	Y				
ISO 1942	2009-12	N	Dentistry - Vocabulary					
ISO 19980	2005-08	N	Ophthalmic instruments - Corneal topographers	N				
ISO 20072	2009-08	N	Aerosol drug delivery device design verification - Requirements and test methods	N				
ISO 20126	2012-01	N	Dentistry - Manual toothbrushes - General requirements and test methods					
ISO 20127	2005-03	N	Dentistry - Powered toothbrushes - General requirements and test methods					
ISO 20160	2006-05	N	Implants for surgery - Metallic materials - Classification of microstructures for alpha+beta titanium alloy bars	N				
ISO 20301	2006-11	N	Health informatics - Health cards - General characteristics					
ISO 20302	2006-12	N	Health informatics - Health cards - Numbering system and registration procedure for issuer identifiers					

ISO 20776-1	2006-11	N	Clinical laboratory testing and in vitro diagnostic test systems_ - Susceptibility testing of infectious agents and evaluation of performance of antimicrobial susceptibility test devices_ - Part_1: Reference method for testing the in vitro activity of antimicrobial agents against rapidly growing aerobic bacteria involved in infectious diseases	Y				
ISO 20776-2	2007-07	N	Clinical laboratory testing and in vitro diagnostic test systems_ - Susceptibility testing of infectious agents and evaluation of performance of antimicrobial susceptibility test devices_ - Part_2: Evaluation of performance of antimicrobial susceptibility test devices	N				
ISO 20795-1	2008-08	N	Dentistry_ - Base polymers_ - Part_1: Denture base polymers					
ISO 20795-1 Technical C	2009-02	N	Dentistry_ - Base polymers_ - Part_1: Denture base polymers; Technical Corrigendum_1					
ISO 20795-2	2010-03	N	Dentistry_ - Base polymers_ - Part_2: Orthodontic base polymers					
ISO 20857	2010-08	N	Sterilization of health care products_ - Dry heat_ - Requirements for the development, validation and routine control of a sterilization process for medical devices	N				
ISO 21090	2011-02	N	Health informatics_ - Harmonized data types for information interchange					
ISO 21171	2006-05	N	Medical gloves_ - Determination of removable surface powder	Y				
ISO 21530	2004-06	N	Dentistry_ - Materials used for dental equipment surfaces_ - Determination of resistance to chemical disinfectants					
ISO 21531	2009-02	N	Dentistry_ - Graphical symbols for dental instruments					
ISO 21533	2003-06	N	Dentistry_ - Reusable cartridge syringes intended for intraligamentary injections					
ISO 21533 Technical Co	2009-12	N	Dentistry_ - Reusable cartridge syringes intended for intraligamentary injections; Technical Corrigendum_1					
ISO 21534	2007-10	N	Non-active surgical implants_ - Joint replacement implants_ - Particular requirements	Y				

ISO 21535	2007-10	N	Non-active surgical implants_- Joint replacement implants_- Specific requirements for hip-joint replacement implants	Y				
ISO 21536	2007-10	N	Non-active surgical implants_- Joint replacement implants_- Specific requirements for knee-joint replacement implants	Y				
ISO 21549-1	2004-05	N	Health informatics_- Patient healthcard data_- Part_1: General structure					
ISO 21549-2	2004-05	N	Health informatics_- Patient healthcard data_- Part_2: Common objects					
ISO 21549-3	2004-05	N	Health informatics_- Patient healthcard data_- Part_3: Limited clinical data					
ISO 21549-4	2006-11	N	Health informatics_- Patient healthcard data_- Part_4: Extended clinical data					
ISO 21549-5	2008-04	N	Health informatics_- Patient healthcard data_- Part_5: Identification data					
ISO 21549-6	2008-04	N	Health informatics_- Patient healthcard data_- Part_6: Administrative data					
ISO 21549-7	2007-06	N	Health informatics_- Patient healthcard data_- Part_7: Medication data					
ISO 21549-8	2010-06	N	Health informatics_- Patient healthcard data_- Part_8: Links					
ISO 2157	1992-06	N	Dental rotary instruments; nominal diameters and designation code number					
ISO 21606	2007-06	N	Dentistry_- Elastomeric auxiliaries for use in orthodontics					
ISO 21649	2006-06	N	Needle-free injectors for medical use_- Requirements and test methods	Y				
ISO 21667	2010-12	N	Health informatics_- Health indicators conceptual framework					
ISO 21671	2006-07	N	Dentistry_- Rotary polishers					
ISO 21671 AMD 1	2011-04	N	Dentistry_- Rotary polishers; Amendment_1					
ISO 21672-1	2012-04	N	Dentistry_- Periodontal probes_- Part_1: General requirements					
ISO 21969	2009-10	N	High-pressure flexible connections for use with medical gas systems	Y				
ISO 21987	2009-10	N	Ophthalmic optics_- Mounted spectacle lenses	N				
ISO 22112	2005-11	N	Dentistry_- Artificial teeth for dental prostheses					
ISO 22254	2005-08	N	Dentistry_- Manual toothbrushes_- Resistance of tufted portion to deflection					
ISO 22374	2005-09	N	Dentistry_- Dental handpieces_- Electrical-powered scalers and scaler tips					
ISO 22413	2010-06	N	Transfer sets for pharmaceutical preparations_- Requirements and test methods					

ISO 22442-1	2007-12	N	Medical devices utilizing animal tissues and their derivatives_ - Part_1: Application of risk management	Y				
ISO 22442-2	2007-12	N	Medical devices utilizing animal tissues and their derivatives_ - Part_2: Controls on sourcing, collection and handling	Y				
ISO 22442-3	2007-12	N	Medical devices utilizing animal tissues and their derivatives_ - Part_3: Validation of the elimination and/or inactivation of viruses and transmissible spongiform encephalopathy (TSE) agents	Y				
ISO 22523	2006-10	N	External limb prostheses and external orthoses_ - Requirements and test methods	Y				
ISO 22609	2004-12	N	Clothing for protection against infectious agents_ - Medical face masks_ - Test method for resistance against penetration by synthetic blood (fixed volume, horizontally projected)	N				
ISO 22610	2006-07	N	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	Y				
ISO 22612	2005-03	N	Clothing for protection against infectious agents_ - Test method for resistance to dry microbial penetration	Y				
ISO 22674	2006-11	N	Dentistry_ - Metallic materials for fixed and removable restorations and appliances					
ISO 22675	2006-10	N	Prosthetics_ - Testing of ankle-foot devices and foot units_ - Requirements and test methods	Y				
ISO 22715	2006-04	N	Cosmetics_ - Packaging and labelling					
ISO 22716	2007-11	N	Cosmetics_ - Good Manufacturing Practices (GMP)_ - Guidelines on Good Manufacturing Practices					
ISO 22794	2007-07	N	Dentistry_ - Implantable materials for bone filling and augmentation in oral and maxillofacial surgery_ - Contents of a technical file					
ISO 22803	2004-09	N	Dentistry_ - Membrane materials for guided tissue regeneration in oral and maxillofacial surgery_ - Contents of a technical file					
ISO 22857	2004-04	N	Health informatics_ - Guidelines on data protection to facilitate trans-border flows of personal health information					
ISO 23317	2007-06	N	Implants for surgery_ - In vitro evaluation for apatite-forming ability of implant materials	N				

ISO 23328-1	2003-08	N	Breathing system filters for anaesthetic and respiratory use_ - Part_1: Salt test method to assess filtration performance	Y				
ISO 23328-2	2002-10	N	Breathing system filters for anaesthetic and respiratory use_ - Part_2: Non-filtration aspects	Y				
ISO 23409	2011-02	N	Male condoms_ - Requirements and test methods for condoms made from synthetic materials					
ISO 23500	2011-05	N	Guidance for the preparation and quality management of fluids for haemodialysis and related therapies	N				
ISO 23599	2012-03	N	Assistive products for blind and vision-impaired persons_ - Tactile walking surface indicators					
ISO 23600	2007-11	N	Assistive products for persons with vision impairments and persons with vision and hearing impairments_ - Acoustic and tactile signals for pedestrian traffic lights					
ISO 23640	2011-12	N	In vitro diagnostic medical devices_ - Evaluation of stability of in vitro diagnostic reagents	N				
ISO 23747	2007-07	N	Anaesthetic and respiratory equipment_ - Peak expiratory flow meters for the assessment of pulmonary function in spontaneously breathing humans	Y				
ISO 23908	2011-06	N	Sharps injury protection_ - Requirements and test methods_ - Sharps protection features for single-use hypodermic needles, introducers for catheters and needles used for blood sampling	N				
ISO 24157	2008-07	N	Ophthalmic optics and instruments_ - Reporting aberrations of the human eye	N				
ISO 24214	2006-11	N	Skin barrier for ostomy aids_ - Vocabulary					
ISO 24234	2004-10	N	Dentistry_ - Mercury and alloys for dental amalgam					
ISO 24234 AMD 1	2011-08	N	Dentistry_ - Mercury and alloys for dental amalgam_ - Amendment_1: Requirements for marking and manufacturer's instructions concerning mercury					
ISO 24415-1	2009-04	N	Tips for assistive products for walking_ - Requirements and test methods_ - Part_1: Friction of tips					
ISO 24415-2	2011-08	N	Tips for assistive products for walking_ - Requirements and test methods_ - Part_2: Durability of tips for crutches					

ISO 24500	2010-10	N	Ergonomics_ - Accessible design_ - Auditory signals for consumer products					
ISO 24501	2010-12	N	Ergonomics_ - Accessible design_ - Sound pressure levels of auditory signals for consumer products					
ISO 24502	2010-12	N	Ergonomics_ - Accessible design_ - Specification of age-related luminance contrast for coloured light					
ISO 24503	2011-01	N	Ergonomics_ - Accessible design_ - Tactile dots and bars on consumer products					
ISO 25424	2009-09	N	Sterilization of medical devices_ - Low temperature steam and formaldehyde_ - Requirements for development, validation and routine control of a sterilization process for medical devices	N				
ISO 25539-1	2003-03	N	Cardiovascular implants_ - Endovascular devices_ - Part_1: Endovascular prostheses	Y				
ISO 25539-1 AMD 1	2005-07	N	Cardiovascular implants_ - Endovascular devices_ - Part_1: Endovascular prostheses; Amendment_1: Test methods	Y				
ISO 25539-2	2008-09	N	Cardiovascular implants_ - Endovascular devices_ - Part_2: Vascular stents	Y				
ISO 25539-3	2011-12	N	Cardiovascular implants_ - Endovascular devices_ - Part_3: Vena cava filters	Y if proposed by CEN				
ISO 25720	2009-08	N	Health informatics_ - Genomic Sequence Variation Markup Language (GSVML)					
ISO 25841	2011-07	N	Female condoms_ - Requirements and test methods					
ISO 26722	2009-04	N	Water treatment equipment for haemodialysis applications and related therapies	N				
ISO 26782	2009-07	N	Anaesthetic and respiratory equipment_ - Spirometers intended for the measurement of time forced expired volumes in humans	Y				
ISO 26782 Technical C	2009-11	N	Anaesthetic and respiratory equipment_ - Spirometers intended for the measurement of time forced expired volumes in humans; Technical Corrigendum_1	Y				
ISO 26825	2008-08	N	Anaesthetic and respiratory equipment_ - User-applied labels for syringes containing drugs used during anaesthesia_ - Colours, design and performance	N				

ISO 27020	2010-12	N	Dentistry_- Brackets and tubes for use in orthodontics					
ISO 27185	2012-02	N	Cardiac rhythm management devices_- Symbols to be used with cardiac rhythm management device labels, and information to be supplied_- General requirements	N				
ISO 27186	2010-03	N	Active implantable medical devices_- Four-pole connector system for implantable cardiac rhythm management devices_- Dimensional and test requirements	N				
ISO 27427	2010-03	N	Anaesthetic and respiratory equipment_- Nebulizing systems and components	N				
ISO 27799	2008-07	N	Health informatics_- Information security management in health using ISO/IEC_2702					
ISO 28158	2010-07	N	Dentistry_- Integrated dental floss and handles					
ISO 28319	2010-05	N	Dentistry_- Laser welding					
ISO 28399	2011-01	N	Dentistry_- Products for external tooth bleaching					
ISO 28620	2010-02	N	Medical devices_- Non-electrically driven portable infusion devices	N				
ISO 29701	2010-09	N	Nanotechnologies_- Endotoxin test on nanomaterial samples for in vitro systems_- Limulus amoebocyte lysate (LAL) test	N				
ISO 29781	2008-12	N	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					
ISO 29782	2008-12	N	Prostheses and orthoses_- Factors to be considered when specifying a prosthesis for a person who has had a lower limb amputation					
ISO 29783-1	2008-12	N	Prosthetics and orthotics_- Vocabulary_- Part_1: Normal gait	N				
ISO 29941	2010-12	N	Condoms_- Determination of nitrosamines migrating from natural rubber latex condoms					
ISO 29942	2011-07	N	Prophylactic dams_- Requirements and test methods					
ISO 3107	2011-03	N	Dentistry_- Zinc oxide/eugenol cements and zinc oxide/non-eugenol cements					
ISO 32	1977-05	N	Gas cylinders for medical use; Marking for identification of content	Y				
ISO 3630-1	2008-02	N	Dentistry_- Root-canal instruments_- Part_1: General requirements and test methods					

ISO 3630-2	2000-12	N	Dental root-canal instruments_ - Part_2: Enlargers					
ISO 3630-3	1994-03	N	Dental root-canal instruments; part_3: condensers, pluggers and spreaders					
ISO 3630-4	2009-07	N	Dentistry_ - Root canal instruments_ - Part_4: Auxiliary instruments					
ISO 3630-5	2011-10	N	Dentistry_ - Endodontic instruments_ - Part_5: Shaping and cleaning instruments					
ISO 3823-1	1997-08	N	Dental rotary instruments_ - Burs_ - Part_1: Steel and carbide burs					
ISO 3823-2	2003-05	N	Dentistry_ - Rotary bur instruments_ - Part_2: Finishing burs					
ISO 3823-2 AMD 1	2008-07	N	Dentistry_ - Rotary bur instruments_ - Part_2: Finishing burs; Amendment_1					
ISO 3826-1	2003-11	N	Plastics collapsible containers for human blood and blood components_ - Part_1: Conventional containers	N				
ISO 3826-2	2008-08	N	Plastics collapsible containers for human blood and blood components_ - Part_2: Graphical symbols for use on labels and instruction leaflets	Y				
ISO 3826-3	2006-09	N	Plastics collapsible containers for human blood and blood components_ - Part_3: Blood bag systems with integrated features	Y				
ISO 389-1	1998-11	N	Acoustics_ - Reference zero for the calibration of audiometric equipment_ - Part_1: Reference equivalent threshold sound pressure levels for pure tones and supra-aural earphones	N				
ISO 389-2	1994-07	N	Acoustics_ - Reference zero for the calibration of audiometric equipment_ - Part_2: Reference equivalent threshold sound pressure levels for pure tones and insert earphones	N				
ISO 389-3	1994-10	N	Acoustics_ - Reference zero for the calibration of audiometric equipment_ - Part_3: Reference equivalent threshold force levels for pure tones and bone vibrators	N				
ISO 389-3 Technical C	1995-08	N	Acoustics_ - Reference zero for the calibration of audiometric equipment_ - Part_3: Reference equivalent treshold force levels for pure tones and bone vibrators; Technical corrigendum_1	N				
ISO 389-4	1994-10	N	Acoustics_ - Reference zero for the calibration of audiometric equipment_ - Part_4: Reference levels for narrow-band masking noise	N				

ISO 389-6	2007-07	N	Acoustics_- Reference zero for the calibration of audiometric equipment_- Part_6: Reference threshold of hearing for test signals of short duration	N				
ISO 389-7	2005-11	N	Acoustics_- Reference zero for the calibration of audiometric equipment_- Part_7: Reference threshold of hearing under free-field and diffuse-field listening conditions	N				
ISO 389-8	2004-05	N	Acoustics_- Reference zero for the calibration of audiometric equipment_- Part_8: Reference equivalent threshold sound pressure levels for pure tones and circumaural earphones	N				
ISO 389-9	2009-05	N	Acoustics_- Reference zero for the calibration of audiometric equipment_- Part_9: Preferred test conditions for the determination of reference hearing threshold levels	N				
ISO 3950	2009-05	N	Dentistry_- Designation system for teeth and areas of the oral cavity					
ISO 3964	1982-12	N	Dental handpieces; Coupling dimensions					
ISO 4049	2009-10	N	Dentistry_- Polymer-based restorative materials					
ISO 4073	2009-07	N	Dentistry_- Information system on the location of dental equipment in the working area of the oral health care provider					
ISO 4074	2002-02	N	Natural latex rubber condoms_- Requirements and test methods	Y				
ISO 4074 Technical Corr	2003-10	N	Natural latex rubber condoms_- Requirements and test methods; Technical Corrigendum_1	N				
ISO 4074 Technical Corr	2008-04	N	Natural latex rubber condoms_- Requirements and test methods; Technical Corrigendum_2	N				
ISO 4135	2001-08	N	Anaesthetic and respiratory equipment_- Vocabulary	Y				
ISO 4823	2000-12	N	Dentistry_- Elastomeric impression materials					
ISO 4823 AMD 1	2007-07	N	Dentistry_- Elastomeric impression materials; Amendment_1					
ISO 4823 Technical Corr	2004-07	N	Dentistry_- Elastomeric impression materials; Technical Corrigendum_1					
ISO 5356-1	2004-05	N	Anaesthetic and respiratory equipment_- Conical connectors_- Part_1: Cones and sockets	N				
ISO 5356-2	2006-09	N	Anaesthetic and respiratory equipment_- Conical connectors_- Part_2: Screw-threaded weight-bearing connectors	Y				

ISO 5358	1992-01	N	Anaesthetic machines for use with humans	N				
ISO 5359	2008-06	N	Low-pressure hose assemblies for use with medical gases	Y				
ISO 5359 AMD 1	2011-12	N	Low-pressure hose assemblies for use with medical gases; Amendment_1	Y				
ISO 5360	2012-01	N	Anaesthetic vaporizers_- Agent-specific filling systems	N				
ISO 5361	1999-09	N	Anaesthetic and respiratory equipment_- Tracheal tubes and connectors	N				
ISO 5361-4	1987-12	N	Tracheal tubes; Part 4 : Cole type	N				
ISO 5362	2006-06	N	Anaesthetic reservoir bags	N				
ISO 5364	2008-07	N	Anaesthetic and respiratory equipment_- Oropharyngeal airways	N				
ISO 5366-1	2000-12	N	Anaesthetic and respiratory equipment_- Tracheostomy tubes_- Part_1: Tubes and connectors for use in adults	Y				
ISO 5366-3	2001-08	N	Anaesthetic and respiratory equipment_- Tracheostomy tubes_- Part_3: Paediatric tracheostomy tubes	N				
ISO 5366-3 Technical	2003-01	N	Anaesthetic and respiratory equipment_- Tracheostomy tubes_- Part_3: Paediatric tracheostomy tubes; Technical Corrigendum_1	N				
ISO 5367	2000-06	N	Breathing tubes intended for use with anaesthetic apparatus and ventilators	N				
ISO 5832-1	2007-06	N	Implants for surgery_- Metallic materials_- Part_1: Wrought stainless steel	N				
ISO 5832-1 Technical	2008-04	N	Implants for surgery_- Metallic materials_- Part_1: Wrought stainless steel; Technical Corrigendum_1	N				
ISO 5832-11	1994-09	N	Implants for surgery_- Metallic materials_- Part_11: Wrought titanium 6-aluminium 7-niobium alloy	N				
ISO 5832-12	2007-05	N	Implants for surgery_- Metallic materials_- Part_12: Wrought cobalt-chromium-molybdenum alloy	N				
ISO 5832-12 Technical	2008-09	N	Implants for surgery_- Metallic materials_- Part_12: Wrought cobalt-chromium-molybdenum alloy; Technical Corrigendum_1	N				

ISO 5832-14	2007-10	N	Implants for surgery_ - Metallic materials_ - Part_14: Wrought titanium 15-molybdenum 5-zirconium 3-aluminium alloy	N				
ISO 5832-2	1999-07	N	Implants for surgery_ - Metallic materials_ - Part_2: Unalloyed titanium	N				
ISO 5832-3	1996-07	N	Implants for surgery_ - Metallic materials_ - Part_3: Wrought titanium 6-aluminium 4-vanadium alloy	N				
ISO 5832-4	1996-07	N	Implants for surgery_ - Metallic materials_ - Part_4: Cobalt-chromium-molybdenum casting alloy	N				
ISO 5832-5	2005-10	N	Implants for surgery_ - Metallic materials_ - Part_5: Wrought cobalt-chromium-tungsten-nickel alloy	N				
ISO 5832-6	1997-07	N	Implants for surgery_ - Metallic materials_ - Part_6: Wrought cobalt-nickel-chromium-molybdenum alloy	N				
ISO 5832-7	1994-02	N	Implants for surgery; metallic materials; part_7: forgeable and cold-formed cobalt-chromium-nickel-molybdenum-iron alloy	N				
ISO 5832-8	1997-07	N	Implants for surgery_ - Metallic materials_ - Part_8: Wrought cobalt-nickel-chromium-molybdenum-tungsten-iron alloy	N				
ISO 5832-9	2007-06	N	Implants for surgery_ - Metallic materials_ - Part_9: Wrought high nitrogen stainless steel	N				
ISO 5833	2002-05	N	Implants for surgery_ - Acrylic resin cements	N				
ISO 5834-1	2005-06	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_1: Powder form	N				
ISO 5834-1 Technical	2007-05	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_1: Powder form; Technical Corrigendum_1	N				
ISO 5834-2	2011-08	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_2: Moulded forms	N				
ISO 5834-3	2005-07	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_3: Accelerated ageing methods	N				
ISO 5834-4	2005-05	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_4: Oxidation index measurement method	N				

ISO 5834-5	2005-06	N	Implants for surgery_ - Ultra-high-molecular-weight polyethylene_ - Part_5: Morphology assessment method	N				
ISO 5835	1991-01	N	Implants for surgery; metal bone screws with hexagonal drive connection, spherical under-surface of head, asymmetrical thread; dimensions	N				
ISO 5836	1988-12	N	Implants for surgery; metal bone plates; holes corresponding to screws with asymmetrical thread and spherical under-surface	N				
ISO 5837-1	1985-06	N	Implants for surgery; Intramedullary nailing systems; Part 1 : Intramedullary nails with cloverleaf or V-shaped cross-section	N				
ISO 5837-2	1980-11	N	Implants for surgery; Intramedullary nailing systems; Part 2 : Medullary pins	N				
ISO 5838-1	1995-11	N	Implants for surgery_ - Skeletal pins and wires_ - Part_1: Material and mechanical requirements	N				
ISO 5838-2	1991-01	N	Implants for surgery; skeletal pins and wires; part_2: Steinmann skeletal pins; dimensions	N				
ISO 5838-3	1993-09	N	Implants for surgery; skeletal pins and wires; part_3: Kirschner skeletal wires	N				
ISO 5840	2005-03	N	Cardiovascular implants_ - Cardiac valve prostheses	Y				
ISO 5841-2	2000-10	N	Implants for surgery_ - Cardiac pacemakers_ - Part_2: Reporting of clinical performance of populations of pulse generators or leads					
ISO 5841-3	2000-10	N	Implants for surgery_ - Cardiac pacemakers_ - Part_3: Low-profile connectors [IS-1] for implantable pacemakers	N				
ISO 5841-3 Technical	2003-11	N	Implants for surgery_ - Cardiac pacemakers_ - Part_3: Low-profile connectors (IS-1) for implantable pacemakers; Technical Corrigendum_1	N				
ISO 594-1	1986-06	N	Conical fittings with a 6 % (Luer) taper for syringes, needles and certain other medical equipment; Part 1 : General requirements	Y		EN 20594-1		

ISO 594-2	1998-09	N	Conical fittings with 6%_(Luer) taper for syringes, needles and certain other medical equipment_- Part_2: Lock fittings	N				
ISO 595-1	1986-12	N	Reusable all-glass or metal-and-glass syringes for medical use; Part 1 : Dimensions	N				
ISO 595-2	1987-12	N	Reusable all-glass or metal-and-glass syringes for medical use; Part 2 : Design, performance requirements and tests	N				
ISO 6009	1992-12	N	Hypodermic needles for single use; colour coding for identification	N				
ISO 6009 Technical Co	2008-03	N	Hypodermic needles for single use_- Colour coding for identification; Technical Corrigendum_1	N				
ISO 6360-1	2004-04	N	Dentistry_- Number coding system for rotary instruments_- Part_1: General characteristics					
ISO 6360-1 Technical Co	2007-09	N	Dentistry_- Number coding system for rotary instruments_- Part_1: General characteristics; Technical Corrigendum_1					
ISO 6360-2	2004-11	N	Dentistry_- Number coding system for rotary instruments_- Part_2: Shapes					
ISO 6360-2 AMD 1	2011-12	N	Dentistry_- Number coding system for rotary instruments_- Part_2: Shapes; Amendment_1					
ISO 6360-3	2005-11	N	Dentistry_- Number coding system for rotary instruments_- Part_3: Specific characteristics of burs and cutters					
ISO 6360-4	2004-06	N	Dentistry_- Number coding system for rotary instruments_- Part_4: Specific characteristics of diamond instruments					
ISO 6360-5	2007-12	N	Dentistry_- Number coding system for rotary instruments_- Part_5: Specific characteristics of root-canal instruments					
ISO 6360-6	2004-06	N	Dentistry_- Number coding system for rotary instruments_- Part_6: Specific characteristics of abrasive instruments					
ISO 6360-7	2006-02	N	Dentistry_- Number coding system for rotary instruments_- Part_7: Specific characteristics of mandrels and special instruments					
ISO 6474-1	2010-02	N	Implants for surgery_- Ceramic materials_- Part_1: Ceramic materials based on high purity alumina	N				
ISO 6475	1989-11	N	Implants for surgery; metal bone screws with asymmetrical thread and spherical under-surface; mechanical requirements and test methods	N				
ISO 6710	1995-08	N	Single-use containers for venous blood specimen collection					

ISO 6872	2008-09	N	Dentistry_ - Ceramic materials					
ISO 6873	1998-03	N	Dental gypsum products					
ISO 6874	2005-08	N	Dentistry_ - Polymer-based pit and fissure sealants					
ISO 6875	2011-07	N	Dentistry_ - Patient chair					
ISO 6876	2001-08	N	Dental root canal sealing materials					
ISO 6877	2006-04	N	Dentistry_ - Root-canal obturating points					
ISO 7151	1988-12	N	Surgical instruments; non-cutting, articulated instruments; general requirements and test methods	N				
ISO 7153-1	1991-04	N	Surgical instruments; metallic materials; part_1: stainless steel	N				
ISO 7153-1 AMD 1	1999-03	N	Surgical instruments_ - Metallic materials_ - Part_1: Stainless steel; Amendment_1	N				
ISO 7176-1	1999-10	N	Wheelchairs_ - Part_1: Determination of static stability					
ISO 7176-10	2008-11	N	Wheelchairs_ - Part_10: Determination of obstacle-climbing ability of electrically powered wheelchairs					
ISO 7176-11	1992-05	N	Wheelchairs; part_11: test dummies					
ISO 7176-13	1989-08	N	Wheelchairs; part_13: determination of coefficient of friction of test surfaces					
ISO 7176-14	2008-02	N	Wheelchairs_ - Part_14: Power and control systems for electrically powered wheelchairs and scooters_ - Requirements and test methods					
ISO 7176-15	1996-11	N	Wheelchairs_ - Part_15: Requirements for information disclosure, documentation and labelling					
ISO 7176-16	1997-05	N	Wheelchairs_ - Part_16: Resistance to ignition of upholstered parts_ - Requirements and test methods					
ISO 7176-19	2008-07	N	Wheelchairs_ - Part_19: Wheeled mobility devices for use as seats in motor vehicles					
ISO 7176-2	2001-06	N	Wheelchairs_ - Part_2: Determination of dynamic stability of electric wheelchairs					
ISO 7176-21	2009-04	N	Wheelchairs_ - Part_21: Requirements and test methods for electromagnetic compatibility of electrically powered wheelchairs and scooters, and battery chargers					
ISO 7176-22	2000-05	N	Wheelchairs_ - Part_22: Set-up procedures					
ISO 7176-23	2002-07	N	Wheelchairs_ - Part_23: Requirements and test methods for attendant-operated stair-climbing devices					
ISO 7176-24	2004-10	N	Wheelchairs_ - Part_24: Requirements and test methods for user-operated stair-climbing devices					
ISO 7176-26	2007-04	N	Wheelchairs_ - Part_26: Vocabulary					
ISO 7176-3	2003-04	N	Wheelchairs_ - Part_3: Determination of effectiveness of brakes					

ISO 7176-4	2008-10	N	Wheelchairs_- Part_4: Energy consumption of electric wheelchairs and scooters for determination of theoretical distance range					
ISO 7176-5	2008-06	N	Wheelchairs_- Part_5: Determination of dimensions, mass and manoeuvring space					
ISO 7176-6	2001-10	N	Wheelchairs_- Part_6: Determination of maximum speed, acceleration and deceleration of electric wheelchairs					
ISO 7176-7	1998-05	N	Wheelchairs_- Part_7: Measurement of seating and wheel dimensions					
ISO 7176-8	1998-07	N	Wheelchairs_- Part_8: Requirements and test methods for static, impact and fatigue strengths					
ISO 7176-9	2009-11	N	Wheelchairs_- Part_9: Climatic tests for electric wheelchairs					
ISO 7193	1985-12	N	Wheelchairs; Maximum overall dimensions					
ISO 7197	2006-06	N	Neurosurgical implants_- Sterile, single-use hydrocephalus shunts and components	Y				
ISO 7197 Technical Corrigendum 1	2007-07	N	Neurosurgical implants_- Sterile, single-use hydrocephalus shunts and components; Technical Corrigendum_1	Y				
ISO 7198	1998-08	N	Cardiovascular implants_- Tubular vascular prostheses	N				
ISO 7199	2009-04	N	Cardiovascular implants and artificial organs_- Blood-gas exchangers (oxygenators)	N				
ISO 7199 AMD 1	2012-02	N	Cardiovascular implants and artificial organs_- Blood-gas exchangers (oxygenators)_- Amendment_1: Clarifications for test methodologies, labelling, and sampling schedule	N				
ISO 7206-1	2008-04	N	Implants for surgery_- Partial and total hip joint prostheses_- Part_1: Classification and designation of dimensions	N				
ISO 7206-10	2003-12	N	Implants for surgery_- Partial and total hip-joint prostheses_- Part_10: Determination of resistance to static load of modular femoral heads	N				
ISO 7206-2	2011-04	N	Implants for surgery_- Partial and total hip joint prostheses_- Part_2: Articulating surfaces made of metallic, ceramic and plastics materials	N				

ISO 7206-4	2010-06	N	Implants for surgery_- Partial and total hip joint prostheses_- Part_4: Determination of endurance properties and performance of stemmed femoral components	N				
ISO 7206-6	1992-03	N	Implants for surgery; partial and total hip joint prostheses; part_6: determination of endurance properties of head and neck region of stemmed femoral components	N				
ISO 7207-1	2007-02	N	Implants for surgery_- Components for partial and total knee joint prostheses_- Part_1: Classification, definitions and designation of dimensions	N				
ISO 7207-2	2011-07	N	Implants for surgery_- Components for partial and total knee joint prostheses_- Part_2: Articulating surfaces made of metal, ceramic and plastics materials	N				
ISO 7376	2009-08	N	Anaesthetic and respiratory equipment_- Laryngoscopes for tracheal intubation	Y				
ISO 7396-1	2007-04	N	Medical gas pipeline systems_- Part_1: Pipeline systems for compressed medical gases and vacuum	Y				
ISO 7396-1 AMD 1	2010-01	N	Medical gas pipeline systems_- Part_1: Pipeline systems for compressed medical gases and vacuum_- Amendment_1: Requirements for terminal units for vacuum fitted on medical supply units with operator-adjustable portions and connected to the pipeline through flexible hoses	Y				
ISO 7396-1 AMD 2	2010-02	N	Medical gas pipeline systems_- Part_1: Pipeline systems for compressed medical gases and vacuum; Amendment_2	Y				
ISO 7396-2	2007-04	N	Medical gas pipeline systems_- Part_2: Anaesthetic gas scavenging disposal systems	Y				
ISO 7405	2008-12	N	Dentistry_- Evaluation of biocompatibility of medical devices used in dentistry	N				
ISO 7439	2011-06	N	Copper-bearing contraceptive intrauterine devices_- Requirements and tests					
ISO 7488	1991-06	N	Dental amalgamators					
ISO 7491	2000-09	N	Dental materials_- Determination of colour stability					
ISO 7492	1997-02	N	Dental explorers					

ISO 7493	2006-05	N	Dentistry_ - Operator's stool					
ISO 7494-1	2011-08	N	Dentistry_ - Dental units_ - Part_1: General requirements and test methods					
ISO 7494-2	2003-03	N	Dentistry_ - Dental units_ - Part_2: Water and air supply					
ISO 7551	1996-12	N	Dental absorbent points					
ISO 7711-1	1997-02	N	Dental rotary instruments_ - Diamond instruments_ - Part_1: Dimensions, requirements, marking and packaging					
ISO 7711-1 AMD 1	2009-05	N	Dental rotary instruments_ - Diamond instruments_ - Part_1: Dimensions, requirements, marking and packaging; Amendment_1					
ISO 7711-2	2011-07	N	Dentistry_ - Rotary diamond instruments_ - Part_2: Discs					
ISO 7711-3	2004-11	N	Dentistry_ - Diamond rotary instruments_ - Part_3: Grit sizes, designation and colour code					
ISO 7740	1985-12	N	Instruments for surgery; Scalpels with detachable blades; Fitting dimensions	N				
ISO 7741	1986-02	N	Instruments for surgery; Scissors and shears; General requirements and test methods	N				
ISO 7785-1	1997-08	N	Dental handpieces_ - Part_1: High-speed air turbine handpieces					
ISO 7785-2	1995-08	N	Dental handpieces_ - Part_2: Straight and geared angle handpieces					
ISO 7786	2001-04	N	Dental rotary instruments_ - Laboratory abrasive instruments					
ISO 7787-1	1984-12	N	Dental rotary instruments; Cutters; Part 1 : Steel laboratory cutters					
ISO 7787-2	2000-12	N	Dental rotary instruments_ - Cutters_ - Part_2: Carbide laboratory cutters					
ISO 7787-3	1991-12	N	Dental rotary instruments; cutters; part_3: carbide laboratory cutters for milling machines					
ISO 7787-4	2002-03	N	Dental rotary instruments_ - Cutters_ - Part_4: Miniature carbide laboratory cutters					
ISO 7864	1993-05	N	Sterile hypodermic needles for single use	N				
ISO 7885	2010-02	N	Dentistry_ - Sterile injection needles for single use	N				
ISO 7886-1	1993-10	N	Sterile hypodermic syringes for single use; part_1: syringes for manual use	N				
ISO 7886-1 Technical	1995-11	N	Sterile hypodermic syringes for single use_ - Part_1: Syringes for manual use; Technical Corrigendum_1	N				
ISO 7886-2	1996-05	N	Sterile hypodermic syringes for single use_ - Part_2: Syringes for use with power-driven syringe pumps	N				

ISO 7886-3	2005-03	N	Sterile hypodermic syringes for single use_- Part_3: Auto-disable syringes for fixed-dose immunization	Y				
ISO 7886-4	2006-10	N	Sterile hypodermic syringes for single use_- Part_4: Syringes with re-use prevention feature	Y				
ISO 7944	1998-06	N	Optics and optical instruments_- Reference wavelengths	N				
ISO 7944 Technical Co	2009-07	N	Optics and optical instruments_- Reference wavelengths; Technical Corrigendum_1	N				
ISO 7998	2005-10	N	Ophthalmic optics_- Spectacle frames_- Lists of equivalent terms and vocabulary	N				
ISO 8009	2004-10	N	Mechanical contraceptives_- Reusable natural and silicone rubber contraceptive diaphragms_- Requirements and tests					
ISO 8009 AMD 1	2012-02	N	Mechanical contraceptives_- Reusable natural and silicone rubber contraceptive diaphragms_- Requirements and tests; Amendment_1					
ISO 80369-1	2010-12	N	Small-bore connectors for liquids and gases in healthcare applications_- Part_1: General requirements	N				
ISO 80601-2-12	2011-04	N	Medical electrical equipment_- Part_2-12: Particular requirements for basic safety and essential performance of critical care ventilators	N				
ISO 80601-2-12 Techn	2011-10	N	Medical electrical equipment_- Part_2-12: Particular requirements for basic safety and essential performance of critical care ventilators; Technical Corrigendum_1	N				
ISO 80601-2-13	2011-08	N	Medical electrical equipment_- Part_2-13: Particular requirements for basic safety and essential performance of an anaesthetic workstation	N				
ISO 80601-2-55	2011-12	N	Medical electrical equipment_- Part_2-55: Particular requirements for the basic safety and essential performance of respiratory gas monitors	N				
ISO 80601-2-56	2009-10	N	Medical electrical equipment_- Part_2-56: Particular requirements for basic safety and essential performance of clinical thermometers for body temperature measurement					

ISO 80601-2-61	2011-04	N	Medical electrical equipment_ - Part_2-61: Particular requirements for basic safety and essential performance of pulse oximeter equipment	N				
ISO 81060-1	2007-12	N	Non-invasive sphygmomanometers_ - Part_1: Requirements and test methods for non-automated measurement type	N				
ISO 81060-2	2009-05	N	Non-invasive sphygmomanometers_ - Part_2: Clinical validation of automated measurement type	N				
ISO 81060-2 Technical	2011-02	N	Non-invasive sphygmomanometers_ - Part_2: Clinical validation of automated measurement type; Technical Corrigendum_1	N				
ISO 8185	2007-07	N	Respiratory tract humidifiers for medical use_ - Particular requirements for respiratory humidification systems	Y				
ISO 8194	1987-06	N	Radiation protection; Clothing for protection against radioactive contamination; Design, selection, testing and use					
ISO 8253-1	2010-11	N	Acoustics_ - Audiometric test methods_ - Part_1: Pure-tone air and bone conduction audiometry	N				
ISO 8253-2	2009-12	N	Acoustics_ - Audiometric test methods_ - Part_2: Sound field audiometry with pure-tone and narrow-band test signals	N				
ISO 8253-3	2012-03	N	Acoustics_ - Audiometric test methods_ - Part_3: Speech audiometry	N				
ISO 8282	1994-10	N	Dental equipment_ - Mercury and alloy mixers and dispensers					
ISO 8319-1	1996-05	N	Orthopaedic instruments_ - Drive connections_ - Part_1: Keys for use with screws with hexagon socket heads	N				
ISO 8319-2	1986-10	N	Orthopaedic instruments; Drive connections; Part 2 : Screwdrivers for single slot head screws, screws with cruciate slot and cross-recessed head screws	N				
ISO 8325	2004-09	N	Dentistry_ - Test methods for rotary instruments					
ISO 8359	1996-12	N	Oxygen concentrators for medical use_ - Safety requirements	Y				
ISO 8362-1	2009-12	N	Injection containers and accessories_ - Part_1: Injection vials made of glass tubing	N				

ISO 8362-2	2008-10	N	Injection containers and accessories_ - Part_2: Closures for injection vials	N				
ISO 8362-3	2001-12	N	Injection containers and accessories_ - Part_3: Aluminium caps for injection vials	N				
ISO 8362-4	2011-09	N	Injection containers and accessories_ - Part_4: Injection vials made of moulded glass	N				
ISO 8362-5	2008-10	N	Injection containers and accessories_ - Part_5: Freeze drying closures for injection vials	N				
ISO 8362-6	2010-06	N	Injection containers and accessories_ - Part_6: Caps made of aluminium-plastics combinations for injection vials	N				
ISO 8362-7	2006-04	N	Injection containers and accessories_ - Part_7: Injection caps made of aluminium-plastics combinations without overlapping plastics part	N				
ISO 8429	1986-09	N	Optics and optical instruments; Ophthalmology; Graduated dial scale	N				
ISO 8536-1	2011-09	N	Infusion equipment for medical use_ - Part_1: Infusion glass bottles	N				
ISO 8536-10	2004-10	N	Infusion equipment for medical use_ - Part_10: Accessories for fluid lines for use with pressure infusion equipment	N				
ISO 8536-11	2004-10	N	Infusion equipment for medical use_ - Part_11: Infusion filters for use with pressure infusion equipment	N				
ISO 8536-12	2007-04	N	Infusion equipment for medical use_ - Part_12: Check valves	N				
ISO 8536-2	2010-03	N	Infusion equipment for medical use_ - Part_2: Closures for infusion bottles	N				
ISO 8536-3	2009-06	N	Infusion equipment for medical use_ - Part_3: Aluminium caps for infusion bottles	N				
ISO 8536-4	2010-10	N	Infusion equipment for medical use_ - Part_4: Infusion sets for single use, gravity feed	N				
ISO 8536-5	2004-02	N	Infusion equipment for medical use_ - Part_5: Burette infusion sets for single use, gravity feed	N				
ISO 8536-6	2009-11	N	Infusion equipment for medical use_ - Part_6: Freeze drying closures for infusion bottles	N				

ISO 8536-7	2009-01	N	Infusion equipment for medical use_- Part_7: Caps made of aluminium-plastics combinations for infusion bottles	N				
ISO 8536-8	2004-08	N	Infusion equipment for medical use_- Part_8: Infusion equipment for use with pressure infusion apparatus	N				
ISO 8536-9	2004-10	N	Infusion equipment for medical use_- Part_9: Fluid lines for use with pressure infusion equipment	N				
ISO 8537	2007-10	N	Sterile single-use syringes, with or without needle, for insulin	N				
ISO 8548-1	1989-08	N	Prosthetics and orthotics; limb deficiencies; part_1: method of describing limb deficiencies present at birth	N				
ISO 8548-2	1993-07	N	Prosthetics and orthotics; limb deficiencies; part_2: method of describing lower limb amputation stumps	N				
ISO 8548-3	1993-07	N	Prosthetics and orthotics; limb deficiencies; part_3: method of describing upper limb amputation stumps	N				
ISO 8548-4	1998-07	N	Prosthetics and orthotics_- Limb deficiencies_- Part_4: Description of causal conditions leading to amputation	N				
ISO 8548-5	2003-07	N	Prosthetics and orthotics_- Limb deficiencies_- Part_5: Description of the clinical condition of the person who has had an amputation	N				
ISO 8549-1	1989-07	N	Prosthetics and orthotics; vocabulary; part_1: general terms for external limb protheses and external orthoses					
ISO 8549-2	1989-07	N	Prosthetics and orthotics; vocabulary; part_2: terms relating to external limb protheses and wearers of these protheses					
ISO 8549-3	1989-07	N	Prosthetics and orthotics; vocabulary; part_3: terms relating to external orthoses					
ISO 8551	2003-08	N	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 8596	2009-07	N	Ophthalmic optics_- Visual acuity testing_- Standard optotype and its presentation	N				

ISO 8598	1996-08	N	Optics and optical instruments_ - Focimeters	N				
ISO 8598 Technical C	1998-05	N	Optics and optical instruments_ - Focimeters; Technical corrigendum_1	N				
ISO 8600-1	2005-05	N	Optics and photonics_ - Medical endoscopes and endotherapy devices_ - Part_1: General requirements	N				
ISO 8600-2	2002-08	N	Optics and optical instruments_ - Medical endoscopes and endoscopic accessories_ - Part_2: Particular requirements for rigid bronchoscopes	N				
ISO 8600-3	1997-07	N	Optics and optical instruments_ - Medical endoscopes and endoscopic accessories_ - Part_3: Determination of field of view and direction of view of endoscopes with optics	N				
ISO 8600-3 AMD 1	2003-12	N	Optics and optical instruments_ - Medical endoscopes and endoscopic accessories_ - Part_3: Determination of field of view and direction of view of endoscopes with optics; Amendment_1	N				
ISO 8600-4	1997-07	N	Optics and optical instruments_ - Medical endoscopes and certain accessories_ - Part_4: Determination of maximum width of insertion portion	N				
ISO 8600-5	2005-03	N	Optics and photonics_ - Medical endoscopes and endotherapy devices_ - Part_5: Determination of optical resolution of rigid endoscopes with optics	N				
ISO 8600-6	2005-03	N	Optics and photonics_ - Medical endoscopes and endotherapy devices_ - Part_6: Vocabulary	N				
ISO 8612	2009-10	N	Ophthalmic instruments_ - Tonometers	N				
ISO 8615	1991-11	N	Implants for surgery; fixation devices for use in the ends of the femur in adults	N				
ISO 8624	2011-02	N	Ophthalmic optics_ - Spectacle frames_ - Measuring system and terminology	N				
ISO 8637	2010-07	N	Cardiovascular implants and extracorporeal systems_ - Haemodialysers, haemodiafilters, haemofilters and haemoconcentrators	N				

ISO 8638	2010-07	N	Cardiovascular implants and extracorporeal systems_- Extracorporeal blood circuit for haemodialysers, haemodiafilters and haemofilters	N				
ISO 8669-1	1988-07	N	Urine collection bags; part_1: vocabulary					
ISO 8669-2	1996-12	N	Urine collection bags_- Part_2: Requirements and test methods					
ISO 8670-1	1988-07	N	Ostomy collection bags; part_1: vocabulary					
ISO 8670-2	1996-12	N	Ostomy collection bags_- Part_2: Requirements and test methods					
ISO 8670-3	2000-03	N	Ostomy collection bags_- Part_3: Determination of odour transmission of colostomy and ileostomy bags					
ISO 8827	1988-10	N	Implants for surgery; staples with parallel legs for orthopaedic use; general requirements	N				
ISO 8828	1988-10	N	Implants for surgery; guidance on care and handling of orthopaedic implants	N				
ISO 8835-7	2011-11	N	Inhalational anaesthesia systems_- Part_7: Anaesthetic systems for use in areas with limited logistical supplies of electricity and anaesthetic gases	Y if proposed by CEN				
ISO 8836	2007-09	N	Suction catheters for use in the respiratory tract	N				
ISO 8871-1	2003-10	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_1: Extractables in aqueous autoclavates	N				
ISO 8871-2	2003-10	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_2: Identification and characterization	N				
ISO 8871-2 AMD 1	2005-07	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_2: Identification and characterization; Amendment_1	N				
ISO 8871-3	2003-08	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_3: Determination of released- particle count	N				
ISO 8871-4	2006-06	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_4: Biological requirements and test methods	N				
ISO 8871-5	2005-08	N	Elastomeric parts for parenterals and for devices for pharmaceutical use_- Part_5: Functional requirements and testing	N				

ISO 8872	2003-03	N	Aluminium caps for transfusion, infusion and injection bottles. - General requirements and test methods	N				
ISO 8980-1	2004-02	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_1: Specifications for single-vision and multifocal lenses	N				
ISO 8980-1 Technical	2006-08	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_1: Specifications for single-vision and multifocal lenses; Technical Corrigendum_1	N				
ISO 8980-2	2004-02	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_2: Specifications for progressive power lenses	N				
ISO 8980-2 Technical	2006-08	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_2: Specifications for progressive power lenses; Technical Corrigendum_1	N				
ISO 8980-3	2003-10	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_3: Transmittance specifications and test methods	N				
ISO 8980-4	2006-08	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_4: Specifications and test methods for anti-reflective coatings	N				
ISO 8980-5	2005-08	N	Ophthalmic optics_- Uncut finished spectacle lenses_- Part_5: Minimum requirements for spectacle lens surfaces claimed to be abrasion-resistant	N				
ISO 9168	2009-07	N	Dentistry_- Hose connectors for air driven dental handpieces					
ISO 9170-1	2008-07	N	Terminal units for medical gas pipeline systems_- Part_1: Terminal units for use with compressed medical gases and vacuum	Y				
ISO 9170-2	2008-07	N	Terminal units for medical gas pipeline systems_- Part_2: Terminal units for anaesthetic gas scavenging systems	Y				
ISO 9173-1	2006-06	N	Dentistry_- Extraction forceps_- Part_1: General requirements and test methods					
ISO 9173-2	2010-05	N	Dentistry_- Extraction forceps_- Part_2: Designation					
ISO 9187-1	2010-10	N	Injection equipment for medical use_- Part_1: Ampoules for injectables	N				

ISO 9187-2	2010-10	N	Injection equipment for medical use_- Part_2: One-point-cut (OPC) ampoules	N				
ISO 9268	1988-12	N	Implants for surgery; metal bone screws with conical under-surface of head; dimensions	N				
ISO 9269	1988-12	N	Implants for surgery; metal bone plates; holes and slots corresponding to screws with conical under-surface	N				
ISO 9333	2006-07	N	Dentistry_- Brazing materials					
ISO 9342-1	2005-05	N	Optics and optical instruments_- Test lenses for calibration of focimeters_- Part_1: Test lenses for focimeters used for measuring spectacle lenses	N				
ISO 9342-2	2005-11	N	Optics and optical instruments_- Test lenses for calibration of focimeters_- Part_2: Test lenses for focimeters used for measuring contact lenses	N				
ISO 9360-1	2000-03	N	Anaesthetic and respiratory equipment_- Heat and moisture exchangers (HMEs) for humidifying respired gases in humans_- Part_1: HMEs for use with minimum tidal volumes of 250_ml	Y				
ISO 9360-2	2001-04	N	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	Y				
ISO 9386-1	2000-11	N	Power-operated lifting platforms for persons with impaired mobility_- Rules for safety, dimensions and functional operation_- Part_1: Vertical lifting platforms					
ISO 9386-2	2000-11	N	Power-operated lifting platforms for persons with impaired mobility_- Rules for safety, dimensions and functional operation_- Part_2: Powered stairlifts for seated, standing and wheelchair users moving in an inclined plane					
ISO 9394	1998-08	N	Ophthalmic optics_- Contact lenses and contact lens care products_- Determination of biocompatibility by ocular study with rabbit eyes	N				
ISO 9583	1993-10	N	Implants for surgery; non-destructive testing; liquid penetrant inspection of metallic surgical implants	N				

ISO 9584	1993-10	N	Implants for surgery; non-destructive testing; radiographic examination of cast metallic surgical implants	N				
ISO 9585	1990-12	N	Implants for surgery; determination of bending strength and stiffness of bone plates	N				
ISO 9626	1991-09	N	Stainless steel needle tubing for manufacture of medical devices	N				
ISO 9626 AMD 1	2001-06	N	Stainless steel needle tubing for the manufacture of medical devices; Amendment_1	N				
ISO 9680	2007-06	N	Dentistry_- Operating lights					
ISO 9687	1993-02	N	Dental equipment; graphical symbols					
ISO 9693	1999-12	N	Metal-ceramic dental restorative systems					
ISO 9693 AMD 1	2005-10	N	Metal-ceramic dental restorative systems; Amendment_1					
ISO 9693-1	2012-02	N	Dentistry_- Compatibility testing_- Part_1: Metal-ceramic systems					
ISO 9713	2002-09	N	Neurosurgical implants_- Self-closing intracranial aneurysm clips	Y				
ISO 9714-1	1991-03	N	Orthopaedic drilling instruments; part_1: drill bits, taps and countersink cutters	N				
ISO 9801	2009-12	N	Ophthalmic instruments_- Trial case lenses	N				
ISO 9873	1998-11	N	Dental hand instruments_- Reusable mirrors and handles					
ISO 9873 Technical Corr	2000-06	N	Dental hand instruments_- Reusable mirrors and handles; Technical Corrigendum_1					
ISO 9917-1	2007-10	N	Dentistry_- Water-based cements_- Part_1: Powder/liquid acid-base cements					
ISO 9917-2	2010-04	N	Dentistry_- Water-based cements_- Part_2: Resin-modified cements					
ISO 9949-1	1993-07	N	Urine absorbing aids; vocabulary; part_1: conditions of urinary incontinence					
ISO 9949-2	1993-07	N	Urine absorbing aids; vocabulary; part_2: products					
ISO 9949-3	1993-07	N	Urine absorbing aids; vocabulary; part_3: identification of product types					
ISO 9997	1999-12	N	Dental cartridge syringes					
ISO 9999	2011-07	N	Assistive products for persons with disability_- Classification and terminology					
ISO/HL7 10781	2009-11	N	Electronic Health Record-System Functional Model, Release_1.1					
ISO/HL7 21731	2006-08	N	Health informatics_- HL_7 version_3_- Reference information model_- Release_1					
ISO/HL7 27931	2009-07	N	Data Exchange Standards_- Health Level Seven Version_2.5_- An application protocol for electronic data exchange in healthcare environments					

ISO/HL7 27932	2009-12	N	Data Exchange Standards_ - HL7 Clinical Document Architecture, Release_2					
ISO/HL7 27951	2009-11	N	Health informatics_ - Common terminology services, release_1					
ISO/HL7 27953-1	2011-12	N	Health informatics_ - Individual case safety reports (ICSRs) in pharmacovigilance_ - Part_1: Framework for adverse event reporting					
ISO/HL7 27953-2	2011-12	N	Health informatics_ - Individual case safety reports (ICSRs) in pharmacovigilance_ - Part_2: Human pharmaceutical reporting requirements for ICSR					
ISO/IEC 10779	2008-06	N	Information technology_ - Office equipment accessibility guidelines for elderly persons and persons with disabilities					
ISO/IEC 13066-1	2011-05	N	Information technology_ - Interoperability with assistive technology (AT)_ - Part_1: Requirements and recommendations for interoperability					
ISO/IEC 29136	2012-05	N	Information technology_ - User interfaces_ - Accessibility of personal computer hardware					
ISO/IEC TR 19765	2007-07	N	Information technology_ - Survey of icons and symbols that provide access to functions and facilities to improve the use of information technology products by the elderly and persons with disabilities					
ISO/IEC TR 19766	2007-06	N	Information technology_ - Guidelines for the design of icons and symbols accessible to all users, including the elderly and persons with disabilities					
ISO/IEC TR 29138-1	2009-06	N	Information technology_ - Accessibility considerations for people with disabilities_ - Part_1: User needs summary					
ISO/IEC TR 29138-2	2009-06	N	Information technology_ - Accessibility considerations for people with disabilities_ - Part_2: Standards inventory					
ISO/IEC TR 29138-3	2009-06	N	Information technology_ - Accessibility considerations for people with disabilities_ - Part_3: Guidance on user needs mapping					
ISO/IEEE 11073-1010	2004-12	N	Health informatics_ - Point-of-care medical device communication_ - Part_10101: Nomenclature	N				
ISO/IEEE 11073-1020	2004-12	N	Health informatics_ - Point-of-care medical device communication_ - Part_10201: Domain information model	N				
ISO/IEEE 11073-10404	2010-05	N	Health informatics_ - Personal health device communication_ - Part_10404: Device specialization_ - Pulse oximeter					
ISO/IEEE 11073-10407	2010-05	N	Health informatics_ - Personal health device communication_ - Part_10407: Device specialization_ - Blood pressure monitor					

ISO/IEEE 11073-10408	2010-05	N	Health informatics_ - Point-of-care medical device communication_ - Part_10408: Device specialization_ - Thermometer	N				
ISO/IEEE 11073-10414	2010-05	N	Health informatics_ - Point-of-care medical device communication_ - Part_10415: Device specialization_ - Weighing scale	N				
ISO/IEEE 11073-10417	2010-05	N	Health informatics_ - Personal health device communication_ - Part_10417: Device specialization_ - Glucose meter					
ISO/IEEE 11073-10471	2010-05	N	Health informatics_ - Point-of-care medical device communication_ - Part_10471: Device specialization_ - Independant living activity hub	N				
ISO/IEEE 11073-20101	2004-12	N	Health informatics_ - Point-of care medical device communications_ - Part_20101: Application profiles; Base standard	N				
ISO/IEEE 11073-20601	2010-05	N	Health informatics_ - Point-of-care medical device communication_ - Part_20601: Application profile_ - Optimized exchange protocol	N				
ISO/IEEE 11073-30200	2004-12	N	Health informatics_ - Point-of-care medical device communications_ - Part_30200: Transport profile; Cable connected	N				
ISO/IEEE 11073-30300	2004-12	N	Health informatics_ - Point-of-care medical device communications_ - Part_30300: Transport profile; Infrared wireless	N				
ISO/TR 11175	1993-08	N	Dental implants; guidelines for developing dental implants					
ISO/TR 11487	2008-12	N	Health informatics_ - Clinical stakeholder participation in the work of ISO_TC 215					
ISO/TR 11548-1	2001-12	N	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	2001-12	N	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/TR 11633-1	2009-11	N	Health informatics_ - Information security management for remote maintenance of medical devices and medical information systems_ - Part_1: Requirements and risk analysis	N				

ISO/TR 11633-2	2009-11	N	Health informatics_- Information security management for remote maintenance of medical devices and medical information systems_- Part_2: Implementation of an information security management system (ISMS)	N				
ISO/TR 11636	2009-12	N	Health Informatics_- Dynamic on-demand virtual private network for health information infrastructure					
ISO/TR 11991	1995-07	N	Guidance on airway management during laser surgery of upper airway	N				
ISO/TR 12309	2009-12	N	Health informatics_- Guidelines for terminology development organizations					
ISO/TR 12773-1	2009-06	N	Business requirements for health summary records_- Part_1: Requirements					
ISO/TR 12773-2	2009-06	N	Business requirements for health summary records_- Part_2: Environmental scan					
ISO/TR 13154	2009-04	N	Medical electrical equipment_- Deployment, implementation and operational guidelines for indentifying febrile humans using a screening thermograph	N				
ISO/TR 13570-1	2005-04	N	Wheelchairs_- Part_1: Guidelines for the application of the ISO_7176 series on wheelchairs					
ISO/TR 13668	1998-11	N	Digital coding of oral health and care					
ISO/TR 14283	2004-07	N	Implants for surgery_- Fundamental principles	N				
ISO/TR 14292	2012-03	N	Health informatics_- Personal health records_- Definition, scope and context					
ISO/TR 14569-1	2007-05	N	Dental materials_- Guidance on testing of wear_- Part_1: Wear by toothbrushing					
ISO/TR 14969	2004-10	N	Medical devices_- Quality mangement systems_- Guidance on the application of ISO_13485: 2003	N				
ISO/TR 15300	2001-05	N	Dentistry_- Application of OSI clinical codification to the classification and coding of dental products					
ISO/TR 15599	2002-10	N	Digital codification of dental laboratory procedures					
ISO/TR 15599 Technical	2003-10	N	Digital codification of dental laboratory procedures; Technical Corrigendum_1					
ISO/TR 16056-1	2004-07	N	Health informatics_- Interoperability of telehealth systems and networks_- Part_1: Introduction and definitions					
ISO/TR 16056-2	2004-07	N	Health informatics_- Interoperability of telehealth systems and networks_- Part_2: Real-time systems					

ISO/TR 16142	2006-01	N	Medical devices_- Guidance on the selection of standards in support of recognized essential principles of safety and performance of medical devices	N				
ISO/TR 17119	2005-01	N	Health informatics_- Health informatics profiling framework					
ISO/TR 18112	2006-01	N	Clinical laboratory testing and in vitro diagnostic test systems_- In vitro diagnostic medical devices for professional use_- Summary of regulatory requirements for information supplied by the manufacturer	N				
ISO/TR 18307	2001-12	N	Health informatics_- Interoperability and compatibility in messaging and communication standards_- Key characteristics					
ISO/TR 20514	2005-10	N	Health informatics_- Electronic health record_- Definition, scope and context					
ISO/TR 20824	2007-07	N	Ophthalmic instruments_- Background for light hazard specification in ophthalmic instrument standards	N				
ISO/TR 21089	2004-06	N	Health informatics_- Trusted end-to-end information flows					
ISO/TR 21548	2010-02	N	Health informatics_- Security requirements for archiving of electronic health records_- Guidelines					
ISO/TR 21730	2007-02	N	Health informatics_- Use of mobile wireless communication and computing technology in healthcare facilities_- Recommendations for electromagnetic compatibility (management of unintentional electromagnetic interference) with medical devices	N				
ISO/TR 22221	2006-11	N	Health informatics_- Good principles and practices for a clinical data warehouse					
ISO/TR 22411	2008-09	N	Ergonomics data and guidelines for the application of ISO/IEC_Guide 71 to products and services to address the needs of older persons and persons with disabilities					
ISO/TR 22442-4	2010-12	N	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	Y if proposed by CEN				

