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				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/RE GIONAL REFERENCE NO.		LEASE 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
		1 1 1 1 1 1	Measurement of electroacoustical					
IEC 60118-0	1983	N	characteristics Hearing aids; part_0: measurement of electroacoustical characteristics;					
IEC 60118-0 AMD 1 IEC 60118-1	1994-01 1995-04	N N	amendment_1 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 Hearing aids Part_1: Hearing aids with					
IEC 60118-1 Edition 3.1	1999-01	N	induction pick-up coil input Hearing aids - Part 12: Dimensions of					
IEC 60118-12	1996-09	N	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 speach-like signal Hearing aids. Part 2 : Hearing aids with					
IEC 60118-2	1983	N	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 Electroacoustics Hearing aids Part_4: Induction loop systems for hearing aid					
IEC 60118-4	2006-10	N	purposes Magnetic field strength Hearing aids. Part 5 : Nipples for insert					
IEC 60118-5	1983	Ν	earphones					

			Hearing aids Part_6: Characteristics of			
IEC 60118-6	1999-06	Ν	electrical input circuits for hearing aids			
120 00118-0	1999-00	IN				
			Electroacoustics Hearing aids Part_7:			
			Measurement of performance characteristics			
			of hearing aids for production, supply and			
IEC 60118-7	2005-10	Ν	delivery quality assurance purposes			
	2000 10	11	Electroacoustics Hearing aids Part_8:			
			Methods of measurement of performance			
			characteristics of hearing aids under simulated			
IEC 60118-8	2005-10	N	in situ working conditions			
			Hearing aids. Part 9 : Methods of			
			measurement of characteristics of hearing			
IEC 60118-9	1985	N	aids with bone vibrator output			
			Electroacoustics Simulators of human head			
			and ear Part_4: Occluded-ear simulator for			
			the measurement of earphones coupled to the			
IEC 60318-4	2010-01	N	ear by means of ear inserts			
			Household and similar electrical appliances			
			Safety Part_2-52: Particular requirements			
IEC 60335-2-52	2005-10	N	for oral hygiene appliances			
			Household and similar electrical appliances			
			Safety Part_2-52: Particular requirements			
IEC 60335-2-52 AMD 1	2008-04	N	for oral hygiene appliances; Amendment_1			
			Household and similar electrical appliances			
	0000.07		Safety Part_2-52: Particular requirements			
IEC 60335-2-52 Edition 3.1	2008-07	N	for oral hygiene appliances			
			Medical electrical equipment X-ray tube			
			assemblies for medical diagnosis			
IEC 60336	2005-04	N	Characteristics of focal spots			
			Medical electrical equipment X-ray tube			
			assemblies for medical diagnosis			
			Characteristics of focal spots;			
IEC 60336 Corrigendum 1	2006-05	Ν	Corrigendum_1			
			Determination of the permanent filtration			
IEC 60522	2003-12	N	of X-ray tube assemblies			
120 00022	2000 12					
			High-voltage cable plug and socket			
IEC 60526	1978	N				
160 00020	19/0	IN	connections for medical X-ray equipment			
			High-voltage cable plug and socket			
IEC 60526 Corrigendum 1	2010-04	N	connections for medical X-ray equipment			
			Medical electrical equipment Dose area			
IEC 60580	2003-09	Ν	product meters			

			Medical electrical equipment Part_1:			
			General requirements for basic safety			
IEC 60601-1	2005-12	Ν	and essential performance			
	2003-12		Medical electrical equipment Part_1:			
			General requirements for basic safety			
			and essential performance;			
IEC 60601-1 Corrigendum 1	2006-12	Ν	Corrigendum_1			
	2000-12		Medical electrical equipment Part_1:			
			General requirements for basic safety			
			and essential performance;			
IEC 60601-1 Corrigendum 2	2007-12	Ν	Corrigendum_2			
	2007-12		Medical electrical equipment Part_1:			
			General requirements for basic safety			
IEC 60601-1 Interpretation S	2008-04	Ν	and essential performance			
	2000-04		Medical electrical equipment Part_1:			
			General requirements for basic safety			
			and essential performance			
IEC 60601-1 Interpretation S	2009-01	N	Interpretation sheet 2			
TEC 00001-1 Interpretation 3	2009-01	IN	Interpretation sheet_2			
			Medical electrical equipment Part_1-1:			
			General requirements for safety;			
			Collateral standard: Safety requirements			
IEC 60601-1-1	2000-12	N	for medical electrical systems			
	2000-12	IN	Medical electrical equipment Part_1-10:			
			General requirements for basic safety			
			and essential performance Collateral			
			Standard: Requirements for the			
			development of physiologic closed-loop			
IEC 60601-1-10	2007-11	Ν	controllers			
	2007-11	IN	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			
IEC 60601-1-11	2010-04	N	healthcare environment			
120 00001-1-11	2010-04	IN	Medical electrical equipment Part_1-11:			
			General requirements for basic safety			
			and essential performance Collateral			
			standard: Requirements for medical			
			electrical equipment and medical			
	2011.04	N	electrical systems used in the home			
IEC 60601-1-11 Corrigendu	12011-04	N	healthcare environment			

			Madical destrict and many Datiest			
			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			
IEC 60601-1-11 Technica	I C 2011-04	Ν	Corrigendum_1			
			Medical electrical equipment Part_1-2:			
			General requirements for basic safety			
			and essential performance Collateral			
			standard: Electromagnetic compatibility			
IEC 60601-1-2	2007-03	Ν	Requirements and tests			
			Medical electrical equipment Part_1-2:			
			General requirements for basic safety			
			and essential performance Collateral			
			standard: Electromagnetic compatibility			
IEC 60601-1-2 Interpretat	ion 2010 02	Ν	Requirements and tests			
	1011 2010-03	IN	Medical electrical equipment Part_1-3:			
			General requirements for basic safety and			
			essential performance Collateral standard:			
			Radiation protection in diagnostic X-ray			
IEC 60601-1-3	2008-01	Ν	equipment			
			Medical electrical equipment Part_1:			
			General requirements for safety			
			4Collateral standard: Programmable			
IEC 60601-1-4	1996-05	Ν	electrical medical systems			
			Medical electrical equipment Part_1-4:			
			General requirements for safety			
			Collateral standard: Programmable			
			electrical medical systems;			
IEC 60601-1-4 AMD 1	1999-10	Ν	Amendment_1			
			Medical electrical equipment Part_1-4:			
			General requirements for safety			
			Collateral standard: Programmable			
IEC 60601-1-4 Edition 1.1	2000-04	Ν	electrical medical systems			
			Medical electrical equipment General			
			requirements for basic safety and			
			essential performance Collateral			
IEC 60601-1-6	2010-01	Ν	Standard: Usability			
IEC 00001-1-0	2010-01	IN	Stanuaru. USability			

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			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				
IEC 60601-1-8	2006-10	N	electrical systems				
	2000 10		Medical electrical equipment Part_1-9:				
			General requirements for basic safety				
			and essential performance Collateral				
			Standard: Requirements for				
IEC 60601-1-9	2007-07	Ν	environmentally conscious design				
			Medical electrical equipment Part_2-1:				
			Particular requirements for the basic				
			safety and essential performance of				
			electron accelerators in the range 1_MeV				
IEC 60601-2-1	2009-10	N	to 50_MeV				
			Medical electrical equipment; part_2:				
			particular requirements for the safety of				
IEC 60601-2-10	1987	N	nerve and muscle stimulators				
			Medical electrical equipment Part_2-10:				
			Particular requirements for the safety of				
			nerve and muscle stimulators;				
IEC 60601-2-10 AMD 1	2001-09	N	Amendment_1				
			Medical electrical equipment Part_2-10:				
			Particular requirements for the safety of				
			nerve and muscle stimulators;				
IEC 60601-2-10 AMD 1 Cc	orri 2002-02	N	Amendment_1				
			Medical electrical equipment Part_2:				
150 00004 0 44	4007.00	N	Particular requirements for the safety of				
IEC 60601-2-11	1997-08	N	gamma beam therapy equipment				
			Amendment_1 Medical electrical				
			equipment Part_2-11: Particular				
IEC 60601-2-11 AMD 1	2004-07	N	requirements for the safety of gamma				
TEC 60601-2-11 AMD 1	2004-07	IN	beam therapy equipment Medical electrical equipment Part_2-13:				
			Particular requirements for the safety and				
			essential performance of anaesthetic				
IEC 60601-2-13	2003-05	N	systems				
	2000-00		Medical electrical equipment Part_2-13:		<u> </u>		
			Particular requirements for the safety and				
			essential performance of anaesthetic				
IEC 60601-2-13 AMD 1	2006-05	Ν	systems; Amendment_1				
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			Medical electrical equipment Part_2-13:			ı
			Particular requirements for the safety of			
IEC 60601-2-13 Edition 3.1	2009-08	Ν	anaesthetic systems			
	2000 00		Medical electrical equipment Part_2-16:			
			Particular requirements for basic safety			
			and essential performance of			
			haemodialysis, haemodiafiltration and			
IEC 60601-2-16	2008-04	Ν	haemofiltration equipment			
	2000 01		Medical electrical equipment Part_2-16:			
			Particular requirements for basic safety			
			and essential performance of			
			haemodialysis, haemodiafiltration and			
IEC 60601-2-16 Corrigendu	un 2008-10	Ν	haemofiltration equipment			
~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			Medical electrical equipment Part_2-17:			
			Particular requirements for the safety of			
			automatically-controlled brachytherapy			
IEC 60601-2-17	2005-09	Ν	afterloading equipment			
			Medical electrical equipment Part_2-18:			
			Particular requirements for basic safety			
			and essential performance of endoscopic			
IEC 60601-2-18	2009-08	Ν	equipment			
			Medical electrical equipment Part_2-19:			
			Particular requirements for the basic			
			safety and essential performance of			
IEC 60601-2-19	2009-02	N	infant incubators	 		
			Medical electrical equipment Part_2-19:			
			Particular requirements for the basic			
			safety and essential performance of			
IEC 60601-2-19 Corrigendu	un 2012-02	N	infant incubators; Corrigendum_1			
			Medical electrical equipment Part_2-2:			
			Particular requirements for the basic			
			safety and essential performance of high			
	2000.02	N	frequency surgical equipment and high			
IEC 60601-2-2	2009-02	N	frequency surgical accessories		+ +	
			Medical electrical equipment Part_2-20:			
			Particular requirements for the basic			
			safety and essential performance of			
IEC 60601-2-20	2009-02	Ν	infant transport incubators			
			Medical electrical equipment Part_2-20:			
			Particular requirements for the basic			
			safety and essential performance of			
			infant transport incubators;			
IEC 60601-2-20 Corrigendu	un 2012-02	Ν	Corrigendum_1			

	<u> </u>		Medical electrical equipment Part_2-21:		<u> </u>	I	
			Particular requirements for the basic				
			safety and essential performance of				
IEC 60601-2-21	2009-02	Ν	infant radiant warmers				
120 00001 2 21	2000 02		Medical electrical equipment Part_2-22:				
			Particular requirements for basic safety				
			and essential performance of surgical,				
			cosmetic, therapeutic and diagnostic				
IEC 60601-2-22	2007-05	Ν	laser equipment				
			Medical electrical equipment Part_2-23:				
			Particular requirements for the basic				
			safety and essential performance of				
			transcutaneous partial pressure				
IEC 60601-2-23	2011-02	Ν	monitoring equipment				
			Medical electrical equipment Part_2-24:				
			Particular requirements for the safety of				
IEC 60601-2-24	1998-02	N	infusion pumps and controllers				
			Medical electrical equipment Part_2-25:				
			Particular requirements for basic safety				
			and essential performance of				
IEC 60601-2-25	2011-10	N	electrocardiographs				
			Medical electrical equipment Part_2-26:				
			Particular requirements for the safety of				
IEC 60601-2-26	2003-12	N	electroencephalographs				
			Medical electrical equipment Part_2-27:				
			Particular requirements for the basic				
			safety and essential performance of				
150 00004 0 05			electrocardiographic monitoring				
IEC 60601-2-27	2011-03	N	equipment				
			Madiaal ale striggt annuis maant - Dant - O.O.				
			Medical electrical equipment Part_2-28:				
			Particular requirements for basic safety				
IEC 60601-2-28	2010-03	Ν	and essential performance of X-ray tube assemblies for medical diagnosis				
IEC 60601-2-28	2010-03	IN	assemblies for medical diagnosis				
			Medical electrical equipment Part_2-29:				
			Particular requirements for the basic				
			safety and essential performance of				
IEC 60601-2-29	2008-06	N	radiotherapy simulators				
	2000 00		Medical electrical equipment; part_2:	1	ł – – ł		
			particular requirements for the safety of				
IEC 60601-2-3	1991-06	Ν	short-wave therapy equipment				
	1331-00	IN	onon wave merapy equipment	1	1		

			Medical electrical equipment Part_2:			
			Particular requirements for the safety of			
			short-wave therapy equipment;			
IEC 60601-2-3 AMD 1	1998-09	Ν	Amendment 1			
120 00001-2-3 AMD 1	1330-03		Medical electrical equipment Part_2-31:			
			Particular requirements for basic safety			
			and essential performance of external			
			cardiac pacemakers with internal power			
IEC 60601-2-31	2008-03	Ν	source			
	2000 00		Medical electrical equipment Part_2-31:			
			Particular requirements for basic safety			
			and essential performance of external			
			cardiac pacemakers with internal power			
IEC 60601-2-31 AMD 1	2011-06	Ν	source			
	2011 00		Medical electrical equipment Part_2-31:			
			Particular requirements for basic safety			
			and essential performance of external			
			cardiac pacemakers with internal power			
IEC 60601-2-31 Edition 2.1	2011-09	Ν	source			
120 00001-2-31 Edition 2.1	2011-03		Medical electrical equipment; part_2:			
			particular requirements for the safety of X-			
IEC 60601-2-32	1994-03	N	ray equipment			
120 00001-2-32	1334-03		Medical electrical equipment Part_2-33:			
			Particular requirements for the basic safety			
			and essential performance of magnetic			
IEC 60601-2-33	2010-03	N	resonance equipment for medical diagnosis			
			Medical electrical equipment Part_2-33:			
			Particular requirements for the basic			
			safety and essential performance of			
			magnetic resonance equipment for			
IEC 60601-2-33 Corrigendum 1	2012-03	N	medical diagnosis			
			Medical electrical equipment Part_2-34:			
			Particular requirements for the basic			
			safety and essential performance of			
			invasive blood pressure monitoring			
IEC 60601-2-34	2011-05	N	equipment			
			Medical electrical equipment Part_2:			
			Particular requirements for the safety of			
			equipment for extracorporeally induced			
IEC 60601-2-36	1997-03	N	lithotripsy			
			Medical electrical equipment Part_2-37:			
			Particular requirements for the basic			
			safety and essential performance of			
			ultrasonic medical diagnostic and			
IEC 60601-2-37	2007-08	N	monitoring equipment			

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			Medical electrical equipment Part_2-39:			
			Particular requirements for basic safety			
			and essential performance of peritoneal			
IEC 60601-2-39	2007-11	N	dialysis equipment			
			Medical electrical equipment Part_2-4:			
			Particular requirements for basic safety			
			and essential performance of cardiac			
IEC 60601-2-4	2010-12	N	defibrillators			
1			Medical electrical equipment Part_2-40:			
			Particular requirements for the safety of			
			electromyographs and evoked response			
IEC 60601-2-40	1998-02	N	equipment			
			Medical electrical equipment Part_2-41:			
			Particular requirements for basic safety			
			and essential performance of surgical			
IEC 60601-2-41	2009-08	N	luminaires and luminaires for diagnosis			
			Medical electrical equipment Part_2-43:			
			Particular requirements for basic safety			
			and essential performance of X-ray			
IEC 60601-2-43	2010-03	Ν	equipment for interventional procedures			
120 00001-2-43	2010-03					
			Medical electrical equipment Part_2-44:			
			Particular requirements for the basic			
			safety and essential performance of X-ray			
IEC 60601-2-44	2009-02	Ν	equipment for computed tomography			
			Medical electrical equipment Part_2-44:			
			Particular requirements for the basic			
			safety and essential performance of X-ray			
IEC 60601-2-44 Corriger	ndun 2010-05	N	equipment for computed tomography			
			Medical electrical equipment Part_2-45:			
			Particular requirements for the basic			
			safety and essential performance of			
			mammographic X-ray equipment and			
IEC 60601-2-45	2011-02	N	mammographic stereotactic devices		 	
			Medical electrical equipment Part_2-46:			
			Particular requirements for the basic			
			safety and essential performance of			
IEC 60601-2-46	2010-12	N	operating tables			

			Medical electrical equipment - Part 2-47:	
			Particular requirements for the basic	
			safety and essential performance of	
IEC 60601-2-47	2012-02	Ν	ambulatory electrocardiographic systems	
			Medical electrical equipment Part_2-49:	
			Particular requirements for the basic	
			safety and essential performance of	
			multifunction patient monitoring	
IEC 60601-2-49	2011-02	Ν	equipment	
			Medical electrical equipment Part_2-5:	
			Particular requirements for basic safety	
			and essential performance of ultrasonic	
IEC 60601-2-5	2009-07	N	physiotherapy equipment	
			Madical destrict any instant. Date 0.50	
			Medical electrical equipment Part_2-50: Particular requirements for the basic	
			safety and essential performance of	
IEC 60601-2-50	2009-03	Ν	infant phototherapy equipment	
	2003-03	IN		
			Medical electrical equipment Part_2-50:	
			Particular requirements for the basic	
			safety and essential performance of	
IEC 60601-2-50 Corrigendur	2010-08	Ν	infant phototherapy equipment	
			Medical electrical equipment Part_2-52:	
			Particular requirements for the basic	
			safety and essential performance of	
IEC 60601-2-52	2009-12	N	medical beds	
			Medical electrical equipment Part_2-52:	
			Particular requirements for the basic	
IEC 60601-2-52 Corrigendur	2010.00	Ν	safety and essential performance of medical beds	
IEC 60601-2-52 Comgendur	2010-09	IN		
			Medical electrical equipment Part_2-52:	
			Particular requirements for the basic	
			safety and essential performance of	
IEC 60601-2-52 Technical C	2010-09	Ν	medical beds; Technical Corrigendum_1	
			IEC_60601-2-54, Ed1: Medical	
			electrical equipment Part_2-54:	
			Particular requirements for the basic	
			safety and essential performance of X-ray	
			equipment for radiography and	
IEC 60601-2-54	2009-06	Ν	radioscopy	

			Medical electrical equipment Part_2-54:	
			Particular requirements for the basic	
			safety and essential performance of X-ray	
			equipment for radiography and	
IEC 60601-2-54 Corrigendu	2010-03	Ν	radioscopy	
			Medical electrical equipment Part_2-54:	
			Particular requirements for the basic	
			safety and essential performance of X-ray	
			equipment for radiography and	
IEC 60601-2-54 Corrigendu	2011-06	Ν	radioscopy	
			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	
IEC 60601-2-57	2011-01	Ν	cosmetic/aesthetic use	
			Medical electrical equipment. Part 2:	
			Particular requirements for the safety of	
IEC 60601-2-6	1984	Ν	microwave therapy equipment	
			Medical electrical equipment Part_2-7:	
			Particular requirements for the safety of	
			high-voltage generators of diagnostic X-	
IEC 60601-2-7	1998-02	Ν	ray generators	
			Medical electrical equipment Part 2-8:	
			Particular requirements for the basic	
			safety and essential performance of	
			therapeutic X-ray equipment operating in	
IEC 60601-2-8	2010-11	Ν	the range 10_kV to 1_MV	
			Medical electrical equipment Part_2:	
			Particular requirements for the safety of	
			therapeutic X-ray equipment in the range	
IEC 60601-2-8 AMD 1	1997-08	Ν	10_kV to 1_MV; Amendment_1	
			Medical electrical equipment Part_2-8:	
			Particular requirements for the safety of	
			therapeutic X-ray equipment operating in	
IEC 60601-2-8 Edition 1.1	1999-04	Ν	the range 10_kV to 1_MV	
			Medical electrical equipment Part_3-1:	
			Essential performance requirements for	
			transcutaneous oxygen and carbon	
			dioxide partial pressure monitoring	
IEC 60601-3-1	1996-07	Ν	equipment	
			Electrical and loading characteristics of X-	
			ray tube assemblies for medical	
IEC 60613	2010-01	N		
IEC 60613	2010-01	Ν	diagnosis	

			Diagnostic X-ray imaging equipment		
			Characteristics of general purpose and		
IEC 60627	2001-08	Ν	mammographic anti-scatter grids		
120 00021	2001.00		Electroacoustics - Audiometric		
			equipment Part_1: Equipment for pure-		
IEC 60645-1	2012-02	Ν	tone audiometry		
			Audiometers; part_2: equipment for		
IEC 60645-2	1993-11	Ν	speech audiometry		
			Electroacoustics Audiometric		
			equipment Part_3: Test signals of short		
IEC 60645-3	2007-03	Ν	duration		
			Electroacoustics Audiometric		
			equipment Part_5: Instruments for the		
			measurement of aural acoustic		
IEC 60645-5	2004-11	N	impedance/admittance		
			Electroacoustics Audiometric		
			equipment Part_6: Instruments for the		
IEC 60645-6	2009-04	N	measurement of otoacoustic emissions		
			Electroacoustics Audiometric		
			equipment Part_7: Instruments for the		
			measurement of auditory brainstem		
IEC 60645-7	2009-04	N	responses		
			Medical electrical equipment		
			Characteristics and test conditions of		
150 00700	0005.40		radionuclide imaging devices Anger		
IEC 60789	2005-10	N	type gamma cameras		
			Medical electrical equipment Characteristics and test conditions of		
			radionuclide imaging devices Anger		
IEC 60789 Corrigendum 1	2009-10	Ν	type gamma cameras; Corrigendum_1		
TEC 60789 Comgendum 1	2009-10	IN	type gamma cameras, Comgendum_1		
			Determination of the maximum		
			symmetrical radiation field from a rotating		
IEC 60806	1984	Ν	anode X-ray tube for medical diagnosis		
120 00000	1001		Medical electrical equipment Medical		
			electron accelerators - Functional		
IEC 60976	2007-10	Ν	performance characteristics		
			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		
IEC 61010-2-040	2005-04	Ν	materials		

	T	I	Cofety requirements for electric-I			
			Safety requirements for electrical			
			equipment for measurement, control and			
			laboratory use Part_2-101: Particular			
			requirements for in vitro diagnostic (IVD)			
IEC 61010-2-101	2002-01	N	medical equipment			
			Standard means for the reporting of the			
			acoustic output of medical diagnostic			
IEC 61157	2007-08	N	ultrasonic equipment			
			Standard means for the reporting of the			
			acoustic output of medical diagnostic			
IEC 61157 Corrigendum 1	2008-08	Ν	ultrasonic equipment; Corrigendum_1			
	2000 00		Radiotherapy simulators; functional			
IEC 61168	1993-12	Ν	performance characteristics			
	1000 12		Ultrasonics; dental descaler systems;			
			measurement and declaration of the output			
IEC 61205	1993-12	N	characteristics			
			Radiotherapy equipment coordinates,			
IEC 61217	2011-12	N	movements and scales			
			Evaluation and routine testing in medical			
			imaging departments Part_2-6:			
			Constancy tests Imaging performance			
			of computed tomography X-ray			
IEC 61223-2-6	2006-11	N	equipment			
			Evaluation and routine testing in medical			
			imaging departments Part_3-2:			
			Acceptance tests Imaging performance			
IEC 61223-3-2	2007-07	N	of mammographic X-ray equipment			
			Evaluation and routine testing in medical			
			imaging departments Part_3-4: Acceptance tests Imaging performance of dental X-ray			
IEC 61223-3-4	2000-03	Ν	equipment			
120 01220 0 4	2000 00		Evaluation and routine testing in medical			
			imaging departments Part_3-5:			
			Acceptance tests Imaging performance			
			of computed tomography X-ray			
IEC 61223-3-5	2004-08	N	equipment			
120 01223-3-3	2004-08	IN	Evaluation and routine testing in medical			
			imaging departments Part_3-5:			
		1				
			Acceptance tests Imaging performance			
IEC 61222 2 E Corrigonation	2006.02	N	of computed tomography X-ray			
IEC 61223-3-5 Corrigendum	2000-03	N	equipment; Corrigendum_1			
IEC 01252 Edition 1.1	2002.02		Electroacoustics Specifications for			
IEC 61252 Edition 1.1	2002-03	N	personal sound exposure meters			

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			
			Medical electrical equipment Characteristics of electro-optical X-ray			
IEC 61262-2	1994-07	N	image intensifiers Part_2: Determination of the conversion factor			
			Medical electrical equipment			
			Characteristics of electro-optical X-ray			
			image intensifiers Part_3:			
	1004.07	N	Determination of the luminance			
IEC 61262-3	1994-07	N	distribution and luminance non-uniformity Medical electrical equipment			
			Characteristics of electro-optical X-ray			
			image intensifiers Part_4:			
IEC 61262-4	1994-07	N	Determination of the image distortion			
			Medical electrical equipment			
			Characteristics of electro-optical X-ray image intensifiers Part_5:			
			Determination of the detective quantum			
IEC 61262-5	1994-07	Ν	efficiency			
			Medical electrical equipment			
			Characteristics of electro-optical X-ray			
			image intensifiers Part_6: Determination of the contrast ratio and			
IEC 61262-6	1994-07	Ν	veiling glare index			
			Medical electrical equipment			
			Characteristics of electro-optical X-ray			
IEC 61262-7	1995-09	Ν	image intensifiers Part-7: Determination of the modulation transfer function			
120 01202-1	1990-09	IN				
			Ultrasonics Hand-held probe Doppler			
			foetal heartbeat detectors Performance			
50 04000	1001.10		requirements and methods of			
IEC 61266	1994-12	N	measurement and reporting Medical diagnostic X-ray equipment			
			Radiation conditions for use in the			
IEC 61267	2005-11	Ν	determination of characetristics			
			Medical electrical equipment			
			Radionuclide calibrators Particular			
IEC 61303	1994-09	N	methods for describing performance			

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			Electrical equipment for measurement,				
			control and laboratory use, control and				
			laboratory use EMC requirements				
			Part_2-6: Particular requirements In-				
IEC 61326-2-6	2005-12	N	vitro diagnostic (IVD) medical equipment				
			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;				
IEC 61326-2-6 Corrigendum	2007-09	N	Corrigendum_1				
			Protective devices against diagnostic medical				
			X-radiation Part_1: Determination of				
IEC 61331-1	1994-10	N	attenuation properties of materials				
			Drotactive devices ansist diamastic modified				
IEC 61331-2	1994-10	Ν	Protective devices against diagnostic medical X-radiation Part_2: Protective glass plates				
IEC 61331-2	1994-10	IN	Protective devices against diagnostic medical				
			X-radiation Part_3: Protective clothing and				
IEC 61331-3	1998-11	Ν	protective devices for gonads				
			Ultrasonics Pulse echo scanners				
			Part_1: Techniques for calibrating spatial				
			measurement systems and				
			measurement of system point-spread				
IEC 61391-1	2006-07	Ν	function response				
	2000 01						
			Ultrasonics - Pulse-echo scanners -				
			Part_2: Measurement of maximum depth				
IEC 61391-2	2010-01	Ν	of penetration and local dynamic range				
	2010 01		Electroacoustics Equipment for the				
			measurement of real-ear acoustical				
IEC 61669	2001-01	N	characteristics of hearing aids				
			Medical electrical equipment				
			Dosimeters with ionization chambers				
			and/or semi-conductor detectors as used				
			in X-ray diagnostic imaging;				
IEC 61674 AMD 1	2002-06	N	Amendment_1				
			Radionuclide imaging devices				
			Characteristics and test conditions				
IEC 61675-1	1998-02	N	Part_1: Positron emission tomographs				
		1	Radionuclide imaging devices				
			Characteristics and test conditions -				
			Part_1: Positron emission tomographs;				
IEC 61675-1 AMD 1	2008-04	Ν	Amendment_1				
			Radionuclide imaging devices				
			Characteristics and test conditions -				
IEC 61675-1 Edition 1.1	2008-06	Ν	Part_1: Positron emission tomographs				
120 01070 1 Edition 1.1	2000 00	1 11	r art_1.1 oonton emission temographs		1	1	

			Radionuclide imaging devices			
			Characteristics and test conditions -			
			Part_2: Single photon emission			
IEC 61675-2	1998-01	N	computed tomographs			
IEC 61675-2	1990-01	IN				
			Radionuclide imaging devices			
			Characteristics and test conditions			
	000440		Part_2: Single photon emission			
IEC 61675-2 AMD 1	2004-12	N	computed tomographs; Amendment_1			
			Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_2: Single photon emission			
IEC 61675-2 Edition 1.1	2005-02	N	computed tomographs			
			Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_3: Gamma camera based			
IEC 61675-3	1998-02	N	wholebody imaging systems			
			Medical electrical equipment			
			Dosimetric instruments used for non-			
			invasive measurement of X-ray tube			
IEC 61676	2002-09	N	voltage in diagnostic radiology			
			Medical electrical equipment			
			Dosimetric instruments used for non-			
			invasive measurement of x-ray tube			
			voltage in diagnostic radiology;			
IEC 61676 AMD 1	2008-11	Ν	Amendment_1			
			Medical electrical equipment			
			Dosimetric instruments used for non-			
			invasive measurement of X-ray tube			
IEC 61676 Edition 1.1	2009-01	Ν	voltage in diagnostic radiology			
			Medical electrical equipment Dosimetric			
			instruments used for non-invasive			
150 04005			measurement of X-ray tube voltage in			
IEC 61685	2002-09	N	diagnostic radiology			
			Ultrasonics Physiotherapy systems			
			Field specifications and methods of			
			measurement in the frequency range			
IEC 61689	2007-08	N	0,5_MHz to 5_MHz			
			Ultrasonics Pressure pulse lithotripters			
IEC 61846	1998-04	N	Characteristics of fields			
			Ultrasonics Surgical systems			
			Measurement and declaration of the			
IEC 61847	1998-01	N	basic output characteristics			
			Medical electrical equipment			
			Requirements for the safety of			
IEC 62083	2009-09	N	radiotherapy treatment planning systems	 		

			Medical electrical equipment Characteristics			
			of digital X-ray imaging devices Part_1: Determination of the detective quantum			
IEC 62127.1	2003-10	N	efficiency			
IEC 02127.1	2003-10	IN				
			Medical electrical equipment			
			Characteristics of digital X-ray imaging			
			devices Part_1: Determination of the			
IEC 62220-1	2003-10	N	detective quantum efficiency			
			Medical electrical equipment			
			Characteristics of digital X-ray imaging			
			devices Part_1-2: Determination of the			
			detective quantum efficiency Detectors			
IEC 62220-1-2	2007-06	N	used in mammography			
			Medical electrical equipment -			
			Characteristics of digital X-ray imaging			
			devices Part_1-3: Determination of the			
			detective quantum efficiency Detectors			
IEC 62220-1-3	2008-06	N	used in dynamic imaging			
120 02220-1-5	2000-00	IN	used in dynamic imaging			
			Medical electrical equipment - Safety of			
150 00074	2005-05	N				
IEC 62274	2005-05	N	radiotheraphy record and verify systems			
			Medical device software Software life			
IEC 62304	2006-05	N	cycle processes			
			Medical electrical equipment Recurrent			
			test and test after repair of medical			
IEC 62353	2007-05	N	electrical equipment			
			Ultrasonics Field characterization			
			Test methods for the determination of			
			thermal and mechanical indices related			
IEC 62359	2010-10	N	to medical diagnostic ultrasonic fields			
			Ultrasonics Field characterization			
			Test methods for the determination of			
			thermal and mechanical indices related			
IEC 62359 Corrigendum 1	2011-03	N	to medical diagnostic ultrasonic fields			
			Medical devices Application of usability			
IEC 62366	2007-10	N	engineering to medical devices			
			Magnetic resonance equipment for medical			
150 00 10 1 1			imaging Part_1: Determination of essential			
IEC 62464-1	2007-01	N	image quality parameters			
			Magnetic resonance equipment for medical			
IEC 62464-2	2010-11	N	imaging Part_2: Classification criteria for pulse sequences			
IEC 02404-2	2010-11	IN IN	puise sequences			

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			Electroacoustics Audio-frequency induction					
			loop systems for assisted hearing Part_1:					
150 63480 4	2010.01	N	Methods of measuring and specifying the performance of system components					
IEC 62489-1	2010-01	N	performance of system components					
			Electroacoustics Audio-frequency induction					
			loop systems for assisted hearing Part_2:					
			Methods of calculating and measuring the low-					
			frequency magnetic field emissions from the					
IEC 62489-2	2011-01	Ν	loop for assessing conformity with guidelines on limits for human exposure					
IEC 02489-2	2011-01	IN	Medical electrical equipment Exposure					
			index of digital X-ray imaging systems					
			Part_1: Definition and requirements of					
IEC 62494-1	2008-08	N	general radiography					
	2000 00		Medical electrical equipment Medical					
			image display systems Part_1:					
IEC 62563-1	2009-12	N	Evaluation methods					
			Application of risk management for IT-					
			networks incorporating medical devices					
			Part_1: Roles, responsibilities and					
IEC 80001-1	2010-10	N	activities					
			Medical electrical equipment Part_2-30:					
			Particular requirements for basic safety					
IEC 80601-2-30	2009-01	N	and essential performance of automated non-invasive sphygnomanometers					
IEC 80601-2-30	2009-01	IN	Medical electrical equipment - Part 2-30:					
			Particular requirements for basic safety					
			and essential performance of automated					
			non-invasive sphygnomanometers;					
IEC 80601-2-30 Corrigend	un 2010-01	N	Corrigendum_1					
			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					
IEC 80601-2-35	2009-10	N	medical use					
			Medical electrical equipment Part_2-35:					
			Particular requirements for the basic					
			safety and essential performance of					
			heating devices using blankets, pads and					
		N	mattresses and intended for heating in					
IEC 80601-2-35 Corrigend	un 2012-03	N	medical use	l				

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			Medical electrical equipment Part_2-58:				
			Particular requirements for basic safety				
			and essential performance of lens				
			removal devices and vitrectomy devices				
IEC 80601-2-58	2008-10	N	for ophthalmic surgery			-	
			Medical electrical equipment Part_2-59:				
			Particular requirements for basic safety				
			and essential performance of screening				
			thermographs for human febrile				
IEC 80601-2-59	2008-10	N	temperature screening				
			Medical electrical equipment Part_2-59:				
			Particular requirements for basic safety				
			and essential performance of screening				
			thermographs for human febrile				
IEC 80601-2-59 Corrigendu	2009-04	Ν	temperature screening; Corrigendum_1				
120 00001-2-35 Comgenad	12003-04	IN	Medical electrical equipment Part_2-60:				
			Particular requirements for basic safety and				
IEC 80601-2-60	2012-02	N	essential performance of dental equipment				
			Medical electrical equipment Glossary				
IEC/TR 60788	2004-02	N	of defined terms				
			Safety of laser products Part_8: Guidelines				
IEC/TR 60825-8	2006-12	Ν	for the safe use of laser beams on humans				
	2000 12		Methods of measuring the performance				
			of ultrasonic pulse-echo diagnostic				
IEC/TR 60854	1986	N	equipment				
			Graphical symbols for electrical				
IEC/TR 60878	2003-07	N	equipment in medical practice				
	2000 01						
			Guidelines for administrative, medical,				
			and nursing staff concerned with the safe				
			use of medical electrical equipment and				
IEC/TR 60930	2008-09	N	medical electrical systems				
			Medical electrical equipment - Medical				
			electron accelerators - Guidelines for				
IEC/TR 60977	2008-07	N	functional performance characteristics				
			Guidelines for the development and use		1	1	1
			of medical electrical equipment				
IEC/TR 61258	2008-08	Ν	educational materials				
			High frequency surgical equipment				
IEC/TR 61289	2011-11	N	Operation and maintenance				
			Nuclear medicine instrumentation -			1	
			Routine tests Part_2: Scintillation				
1			cameras and single photon emission				
IEC/TR 61948-2	2001-02	N	computed tomography imaging				
0, 01010 L			compared tomography imaging		1		

			Nuclear medicine instrumentation				
			Routine tests Part_3: Positron emission				
EC/TR 61948-3	2005-07	Ν	tomographs				
			Nuclear medicine instrumentation				
			Routine tests Part_4: Radionuclide				
IEC/TR 61948-4	2006-11	Ν	calibrators				
			Medical electrical equipment				
			Guidelines for implementation of DICOM				
IEC/TR 62266	2002-03	Ν	in radiotherapy				
20,110 02200	2002 00		Considerations of unaddressed safety				
			aspects in the second edition of				ł
			IEC_60601-1 and proposals for new				
IEC/TR 62296	2009-01	N	requirements				
LO/11 02290	2009-01	IN	Mapping between the clauses of the third				
							I
	0000.05	NI	edition of IEC_60601-1 and the 1988				
EC/TR 62348	2006-05	N	edition as amended				
			General testing procedures for medical				
EC/TR 62354	2009-10	N	electrical equipment				
			Requirements for measurement				
			standards for high intensity therapeutic				
EC/TR 62649	2010-04	N	ultrasound (HITU) devices				
			Audio, video and multimedia systems and				
			equipment Activities and considerations				
EC/TR 62678	2010-10	N	related to accessibility and usability		_		
			Medical device software Part_1: Guidance				
EC/TR 80002-1	2009-09	Ν	on the application of ISO_14971 to medical device software				
EC/TR 80002-1	2009-09	IN	Radiotherapy simulators; guidelines for				
	1993-12	N					
EC/TR2 61170	1993-12	IN	functional performance characteristics				
			Evaluation and routine testing in medical				
			imaging departments; part_1: general				
IEC/TR2 61223-1	1993-07	N	aspects				
			Ultrasonics Real-time pulse-echo				
			systems Test procedures to determine				
IEC/TR2 61390	1996-07	N	the performance specifications				
			Fundamental aspects of safety standards				
IEC/TR3 60513	1994-01	Ν	for medical electrical equipment				
			Cardiac defibrillators; cardiac				
IEC/TR3 61288-1	1993-10	Ν	defibrillators-monitors; part_1: operation				
			Cardiac defibrillators; cardiac				
			defibrillators-monitors; part_2:				
IEC/TR3 61288-2	1993-10	Ν	maintenance				
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			Medical electrical equipment Digital		
			Medical electrical equipment Digital imaging and communications in medicine		
IEC/TR3 61852	1998-04	N	(DICOM) Radiotherapy objects		
120/110301032	1330-04		Guidelines for radiotherapy treatment rooms		
IEC/TR3 61859	1997-04	Ν	design		
			Medical suction equipment Part_1:		
			Electrically powered suction equipment		
ISO 10079-1	1999-08	Ν	Safety requirements		
			Medical suction equipment Part_2:		
ISO 10079-2	1999-08	N	Manually powered suction equipment		
			Medical suction equipment Part_3:		
			Suction equipment powered from a		
ISO 10079-3	1999-08	N	vacuum or pressure source		
			Oxygen concentrator supply systems for		
ISO 10083	2006-07	N	use with medical gas pipeline systems		
			Dentistry Soft lining materials for removable		
ISO 10139-1	2005-02	Ν	dentures Part_1: Materials for short-term use		
130 10139-1	2003-02	IN	Dentistry Soft lining materials for removable		
			dentures Part_1: Materials for short-term		
ISO 10139-1 Technical C	Corrigen 2006-03	Ν	use; Technical Corrigendum_1		
100 10100 0			Dentistry Soft lining materials for removable		
ISO 10139-2	2009-08	N	dentures Part_2: Materials for long-term use Health informatics Messages and	 	
			communication - Web access reference		
ISO 10159	2011-12	Ν	manifest		
			Dentistry Corrosion test methods for metallic		
ISO 10271	2011-08	N	materials		
			Single-use sterile rubber surgical gloves		
ISO 10282	2002-09	N	Specification		
			Single-use sterile rubber surgical gloves		
ISO 10282 Technical	Corrige 2005-06	N	Specification; Technical Corrigendum_1		
			Ophthalmic optics Semi-finished		
			spectacle lens blanks Part_1:		
100 10000 1			Specifications for single-vision and		
ISO 10322-1	2006-02	N	multifocal lens blanks	 	
			Ophthalmic optics Semi-finished		
			spectacle lens blanks Part_2:		
ISO 10322-2	2006-02	Ν	Specifications for progressive power lens blanks		
130 10322-2	2000-02	IN	Dental rotary instruments; bore diameters for		
ISO 10323	1991-11	Ν	discs and wheels		
			Prosthetics Structural testing of lower-limb		
ISO 10328	2006-10	N	prostheses Requirements and test methods		

<b></b>			Implants for surgery Malleable wires for		
100 1000 1	1001.00		use as sutures and other surgical		
ISO 10334	1994-08	N	applications	 	
			Ophthalmic instruments Refractor		
ISO 10341	2009-07	N	heads	 	
			Ophthalmic instruments Eye		
ISO 10342	2010-06	N	refractometers		
			Ophthalmic instruments		
ISO 10343	2009-07	N	Ophthalmometers		
			Dentistry Contents of technical file for dental		
ISO 10451	2010-06	N	implant systems		
100 10177	0004.40		Dentistry Polymer-based crown and bridge		
ISO 10477	2004-10	N	materials		
			Pressure regulators for use with medical		
			gases Part_1: Pressure regulators and		
			pressure regulators with flow-metering		
ISO 10524-1	2006-02	N	devices		
			Pressure regulators for use with medical		
			gases Part_2: Manifold and line		
ISO 10524-2	2005-05	N	pressure regulators	 	
			Pressure regulators for use with medical		
			gases Part_3: Pressure regulators		
ISO 10524-3	2005-05	Ν	integrated with cylinder valves		
			Pressure regulators for use with medical		
ISO 10524-4	2008-06	N	gases Part_4: Low-pressure regulators		
			Hoists for the transfer of disabled		
			persons Requirements and test		
ISO 10535	2006-12	N	methods		
			Technical systems and aids for disabled or		
			handicapped persons Wheelchair tiedown		
			and occupant-restraint systems Part_1:		
			Requirements and test methods for all		
ISO 10542-1	2001-07	N	systems		
			Technical systems and aids for disabled or		
			handicapped persons Wheelchair tiedown and occupant-restraint systems Part_2: Four-		
ISO 10542-2	2001-07	Ν	point strap-type tiedown systems		
130 10342-2	2001-07	IN	Technical systems and aids for disabled or		
			handicapped persons Wheelchair tiedown		
			and occupant-restraint systems Part_3:		
ISO 10542-3	2005-02	Ν	Docking-type tiedown systems		
			Technical systems and aids for disabled or		1
			handicapped persons Wheelchair tiedown		
			and occupant-restraint systems Part_4:		
ISO 10542-4	2004-09	Ν	Clamp-type tiedown systems		

			Technical systems and aids for disabled or			
			handicapped persons Wheelchair tiedown			
ISO 10542 5	2004.04	N	and occupant-restraint systems Part_5:			
ISO 10542-5	2004-04	N	Systems for specific wheelchairs	+		
			Ctarila, single use introvessular			
	1005.00		Sterile, single-use intravascular			
ISO 10555-1	1995-06	N	catheters Part_1: General requirements		-	
			Sterile, single-use intravascular			
			catheters Part_1: General			
ISO 10555-1 AMD 1	1999-07	N	requirements; Amendment_1			
			Sterile, single-use intravascular			
			catheters Part_1: General			
ISO 10555-1 AMD 2	2004-05	N	requirements; Amendment_2			
			Sterile, single-use intravascular			
			catheters Part_2: Angiographic			
ISO 10555-2	1996-06	Ν	catheters			
			Sterile, single-use intravascular			
			catheters Part_2: Angiographic			
ISO 10555-2 Technical Corri	2002-06	Ν	catheters; Technical Corrigendum_1			
	2002 00		Sterile, single-use intravascular			
			catheters Part_3: Central venous			
ISO 10555-3	1996-06	Ν	catheters			
10000-0	1330-00		Sterile, single-use intravascular			
			catheters Part_3: Central venous			
ISO 10555-3 Technical Corri	2002.06	N	catheters; Technical Corrigendum_1			
ISO 10555-5 Technical Com	2002-06	IN	Sterile, single-use intravascular			
	4000.00	N	catheters Part_4: Balloon dilatation			
ISO 10555-4	1996-06	N	catheters			
			Sterile, single-use intravascular			
			catheters Part_4: Balloon dilatation			
ISO 10555-4 Technical Corri	2002-06	N	catheters; Technical Corrigendum_1			
			Sterile, single-use intravascular			
			catheters Part_5: Over-needle			
ISO 10555-5	1996-06	N	peripheral catheters			
			Sterile, single-use intravascular			
			catheters Part_5: Over-needle			
ISO 10555-5 AMD 1	1999-01	Ν	peripheral catheters; Amendment_1			
			Sterile, single-use intravascular			
			catheters Part_5: Over-needle			
			peripheral catheters; Technical			
ISO 10555-5 Technical Corri	2002-06	Ν	Corrigendum 1			
			Dental equipment High- and medium-			
ISO 10637	1999-08	Ν	volume suction systems			
			Dentistry Powered polymerization			
			activators Part_1: Quartz tungsten halogen			
ISO 10650-1	2004-11	Ν	lamps			

			Dentistry Powered polymerization		1
			activators Part_2: Light-emitting diode (LED)		
ISO 10650-2	2007-09	Ν	lamps		
			Lung ventilators for medical use		
			Particular requirements for basic safety		
			and essential performance Part_2:		
			Home care ventilators for ventilator-		
ISO 10651-2	2004-07	N	dependent patients		
			Lung ventilators for medical use		
ISO 10651-3	1997-01	Ν	Part_3: Particular requirements for emergency and transport ventilators		
130 10051-3	1997-01	IN	Lung ventilators Part_4: Particular		
			requirements for operator-powered		
ISO 10651-4	2002-03	Ν	resuscitators		
	2002 00				
			Lung ventilators for medical use		
			Particular requirements for basic safety		
			and essential performance Part_5: Gas-		
ISO 10651-5	2006-02	N	powered emergency resuscitators		
			Lung ventilators for medical use		
			Particular requirements for basic safety and essential performance Part_6:		
ISO 10651-6	2004-07	Ν	Home-care ventilatory support devices		
150 10031-0	2004-07	IN	Ophthalmic optics - Spectacle frames		
			and sunglasses electronic catalogue and		
			identification Part_1: Product		
			identification and electronic catalogue		
ISO 10685-1	2011-12	Ν	product hierarchy		
ISO 10873	2010-09	N	Dentistry Denture adhesives		
			Optics and optical instruments		
			Operation microscopes Part_1:		
ISO 10936-1	2000-06	N	Requirements and test methods		
			Optics and photonics Operation		
			microscopes Part_2: Light hazard from operation microscopes used in ocular		
ISO 10936-2	2010-01	Ν	surgery		
130 10930-2	2010-01	IN	Ophthalmic instruments Chart		
ISO 10938	1998-05	Ν	projectors		
	1000 00		Ophthalmic instruments Slit-lamp		
ISO 10939	2007-02	Ν	microscopes		
			Ophthalmic instruments Fundus		
ISO 10940	2009-08	N	cameras		
			Ophthalmic instruments Direct		
ISO 10942	2006-06	N	ophthalmoscopes		
			Ophthalmic instruments Indirect		
ISO 10943	2011-08	N	ophthalmoscopes		

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ISO 10944	2009-08	N	Ophthalmic instruments Synoptophores				
100 10044	2000 00	11	Caps made of aluminium-plastics				
			combinations for infusion bottles and				
			injection vials Requirements and test				
ISO 10985	2009-02	Ν	methods				
130 10905	2009-02	IN	memous				
			Biological evaluation of medical devices				
			Part_1: Evaluation and testing within a				
ISO 10993-1	2009-10	Ν	risk management process	Y	Ν		
100 10333-1	2003-10	IN .	Biological evaluation of medical devices	I			
			Part 1: Evaluation and testing within a				
			_ 0				
ISO 10993-1 Technica	0.00	N	risk management process; Technical	Y	N		
150 10993-1 Technica	ai Com 2010-06	N	Corrigendum_1 Biological evaluation of medical devices	Ŷ	IN		
100 40000 40	0040.00		Part_10: Tests for irritation and skin				
ISO 10993-10	2010-08	Ν	sensitization	Y	N		
			Biological evaluation of medical devices				
ISO 10993-11	2006-08	Ν		Y	N		
120 10992-11	2000-00	IN	Part_11: Tests for systemic toxicity Biological evaluation of medical devices	ř			
100 40000 40	0007.44	NI	Part_12: Sample preparation and	Ň			
ISO 10993-12	2007-11	N	reference materials	Y	N		
			Biological evaluation of medical devices				
			Part_13: Identification and quantification				
			of degradation products from polymeric				
ISO 10993-13	2010-06	N	medical devices	Y	N		
			Dielegies evolution of medical devices				
			Biological evaluation of medical devices				
100 40000 44	0004.44		Part_14: Identification and quantification				
ISO 10993-14	2001-11	N	of degradation products from ceramics	Y	N		
			Biological evaluation of medical devices				
			Part_15: Identification and quantification				
			of degradation products from metals and				
ISO 10993-15	2000-12	N	alloys	Y	N		
			Distantiant evention of modical devices				
			Biological evaluation of medical devices				
			Part_16: Toxicokinetic study design for				
ISO 10993-16	2010-02	N	degradation products and leachables	Y	N		
			Dislogical evolution of medical devices				
			Biological evaluation of medical devices				
100 40000 17	0000 10	• •	Part_17: Establishment of allowable				
ISO 10993-17	2002-12	N	limits for leachable substances	Y	N		+
			Biological evaluation of medical devices				
			Part_18: Chemical characterization of				
ISO 10993-18	2005-07	N	materials	Y	N		

ISO 10993-2	2006-07	N	Biological evaluation of medical devices Part_2: Animal welfare requirements				
			Biological evaluation of medical devices Part_3: Tests for genotoxicity,				
ISO 10993-3	2003-10	N	carcinogenicity and reproductive toxicity	Y	N	 	
ISO 10993-4	2002-10	N	Biological evaluation of medical devices Part_4: Selection of test for interactions with blood	Y	N		
			Biological evaluation of medical devices Part_4: Selection of tests for interactions				
ISO 10993-4 AMD 1	2006-07	N	with blood	Y	N		
ISO 10993-5	2009-06	N	Biological evaluation of medical devices Part_5: Tests for in vitro cytotoxicity	Y	N		
180 40002 6	2007.04	N	Biological evaluation of medical devices Part_6: Tests for local effects after	V	N		
ISO 10993-6	2007-04	N	implantation Biological evaluation of medical devices	Y	N		
ISO 10993-7	2008-10	N	Part_7: Ethylene oxide sterilization residuals	Y	N		
			Biological evaluation of medical devices Part_7: Ethylene oxide sterilization				
ISO 10993-7 Technical Co	orri 2009-11	Ν	residuals; Technical Corrigendum_1	Y	N		
ISO 10993-9	2009-12	N	Biological evaluation of medical devices Part_9: Framework for identification and quantification of potential degradation products	Y	N		
ISO 11040-1	1992-11	N	Prefilled syringes; part_1: glass cylinders for dental local anaesthetic cartridges				
ISO 11040-2	2011-04	N	Prefilled syringes Part_2: Plunger stoppers for dental local anaesthetic cartridges				
ISO 11040-3	2012-01	Ν	Prefilled syringes Part_3: Seals for dental local anaesthetic cartridges				
ISO 11040-4	2007-02	N	Prefilled syringes Part_4: Glass barrels for injectables				
ISO 11040-5	2012-01	N	Prefilled syringes Part_5: Plunger stoppers for injectables				
ISO 11070	1998-05	N	Sterile single-use intravascular catheter introducers				
ISO 11072 00404	2008.01	N	Health informatics Point-of-care medical device communication Part_90101:				
ISO 11073-90101	2008-01	N	Analytical instruments Point-of-care test				

[			Health informatics Standard communication				
			protocol Part_91064: Computer-assisted				
ISO 11073-91064	2009-05	Ν	electrocardiography				
			Sterilization of health care products -				
			Ethylene oxide Part_1: Requirements				
			for development, validation and routine				
			control of a sterilization process for				
ISO 11135-1	2007-05	Ν	medical devices	Y	N		
			Sterilization of health care products				
			Radiation Part_1: Requirements for				
			development, validation and routine				
			control of a sterilization process for				
ISO 11137-1	2006-04	Ν	medical devices	Y	N	/NZS ISO 11137-1	
			Sterilization of health care products				
			Radiation Part_2: Establishing the				
ISO 11137-2	2012-03	Ν	sterilization dose	Y	N	/NZS ISO 11137-2	
			Sterilization of health care products -				
			Radiation Part_3: Guidance on				
ISO 11137-3	2006-04	Ν	dosimetric aspects	Y	Ν	/NZS ISO 11137-3	
			Sterilization of health care products -				
			Biological indicators Part_1: General				
ISO 11138-1	2006-07	Ν	requirements				
			Sterilization of health care products				
			Biological indicators Part_2: Biological				
			indicators for ethylene oxide sterilization				
ISO 11138-2	2006-07	Ν	processes				
			Sterilization of health care products				
			Biological indicators Part_3: Biological				
			indicators for moist heat sterilization				
ISO 11138-3	2006-07	Ν	processes				
			Sterilization of health care products				
			Biological indicators Part_4: Biological				
			indicators for dry heat sterilization				
ISO 11138-4	2006-07	Ν	processes				
			Sterilization of health care products				
			Biological indicators Part_5: Biological				
			indicators for low-temperature steam and				
ISO 11138-5	2006-07	N	formaldehyde sterilization processes				
			Sterilization of health care products				
			Chemical indicators Part_1: General				
ISO 11140-1	2005-07	N	requirements				
			Sterilization of health care products				
			Chemical indicators Part_3: Class_2				
			indicator systems for use in the Bowie				
ISO 11140-3	2007-03	N	and Dick-type steam penetration test				

<b>F</b>			Starilization of bookb core products	l.	<b>I</b>	
			Sterilization of health care products Chemical indicators - Part 3: Class 2			
			indicator systems for use in the Bowie			
			and Dick-type steam penetration test;			
ISO 11140-3 Technica	al Corri 2007-11	N	Technical Corrigendum_1			
			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			
ISO 11140-4	2007-03	Ν	penetration			
			Sterilization of health care products			
			Chemical indicators Part_5: Class_2			
			indicators for Bowie and Dick-type air			
ISO 11140-5	2007-03	Ν	removal tests			
ISO 11143	2008-07	Ν	Dentistry Amalgam separators			
			Dental equipment Connections for supply			
ISO 11144	1995-05	Ν	and waste lines			
			Packaging Accessible design General			
ISO 11156	2011-07	N	requirements			
			Single-use medical examination gloves			
			Part_1: Specification for gloves made			
ISO 11193-1	2008-09	Ν	from rubber latex or rubber solution			
			Single-use medical examination gloves			
			Part_2: Specification for gloves made			
ISO 11193-2	2006-11	Ν	from poly(vinyl chloride)			
			Gas mixers for medical use - Stand-			
ISO 11195	1995-10	Ν	alone gas mixers			
ISO 11197	2004-12	N	Medical supply units			
	200112		Walking aids manipulated by both arms			
			Requirements and test methods Part_1:			
ISO 11199-1	1999-08	Ν	Walking frames			
			Walking aids manipulated by both arms			
			Requirements and test methods Part_2:			
ISO 11199-2	2005-04	N	Rollators			
			Walking aids manipulated by both arms			
100 11100 0			Requirements and test methods Part_3:			
ISO 11199-3	2005-04	N	Walking tables			
			Cardiac defibrillators Connector			
			assembly DF-1 for implantable			
			defibrillators Dimensional and test			
ISO 11318	2002-08	N	requirements			
			Assistive products for walking manipulated by			
100 44004 4	0007.00	N	one arm Requirements and test methods			
ISO 11334-1	2007-02	N	Part_1: Elbow crutches			
			Walking aids manipulated by one arm Requirements and test methods Part_4:			
ISO 11334-4	1999-02	Ν	Walking sticks with three or more legs			
130 11334-4	1999-02	IN	waiking sucks with three of more legs			

			Transfusion equipment for medical use;				
ISO 1135-3	1986-11	Ν	Part 3 : Blood-taking set				
130 1135-3	1900-11	IN	Fait 5. blood-taking set				
			Transferring a submany for modeling land				
100 1107 1			Transfusion equipment for medical use				
ISO 1135-4	2012-03	N	Part_4: Transfusion sets for single use				
			Optics and optical instruments				
ISO 11380	1994-10	N	Ophthalmic optics Formers				
			Optics and optical instruments				
ISO 11381	1994-12	N	Ophthalmic optics Screw threads				
			Containers and accessories for				
			pharmaceutical preparations Part_1: Drop-				
ISO 11418-1	2005-02	N	dispensing glass bottles				
			Containers and accessories for				
			pharmaceutical preparations Part_2: Screw-				
ISO 11418-2	2005-02	N	neck glass bottles for syrups			 _	-
			Containers and accessories for				
			pharmaceutical preparations Part_3: Screw- neck glass bottles (veral) for solid and liquid				
ISO 11418-3	2005-02	N	dosage forms				
130 11410-3	2003-02	IN	Containers and accessories for				
			pharmaceutical preparations Part_4: Tablet				
ISO 11418-4	2005-02	Ν	glass bottles				
	2000 02		Containers and accessories for				
			pharmaceutical preparations - Part 5:				
ISO 11418-5	1997-12	Ν	Dropper assemblies				
			Containers and accessories for				
			pharmaceutical preparations Part_7: Screw-				
			neck vials made of glass tubing for liquid				
ISO 11418-7	1998-10	N	dosage forms				
			Dental handpieces Dental low-voltage				
ISO 11498	1997-02	N	electrical motors				
			Dentistry Single-use cartridges for local				
ISO 11499	2007-07	N	anaesthetics				
			Packaging for terminally sterilized				
			medical devices Part_1: Requirements				
			for materials, sterile barrier systems and				
ISO 11607-1	2006-04	Ν	packaging systems	Y	N		
			Packaging for terminally sterilized				
			medical devices Part_2: Validation				
			requirements for forming, sealing and				
ISO 11607-2	2006-04	Ν	assembly processes	Y	N		
	2000 07		Pen-injectors for medical use Part_1:	1			
			Pen-injectors; Requirements and test				
ISO 11608-1	2000-12	Ν	methods				
	2000-12	IN	methods				
			Den injectors for medical use				
100 44000 0	0000.40	NI	Pen-injectors for medical use Part_2:				
ISO 11608-2	2000-12	N	Needles; Requirements and test methods				

			Pen-injectors for medical use Part_3:				
			Finished cartridges; Requirements and				
ISO 11608-3	2000-12	Ν	test methods				
130 11008-3	2000-12	IN	Pen-injectors for medical use Part_4:				
			Requirements and test methods for				
			electronic and electromechanical pen-				
100 11609 1	2006 02	N					
ISO 11608-4	2006-03	N	injectors Dentistry Dentifrices Requirements, test				
ISO 11609	2010-09	Ν	methods and marking				
100 11003	2010 05		Quality of dialysis fluid for haemodialysis				
ISO 11663	2009-04	Ν	and related therapies				
	2000 01		Packaging Tactile warnings of danger				
ISO 11683	1997-10	Ν	Requirements				
			Anaesthetic and respiratory equipment				
ISO 11712	2009-05	Ν	Supralaryngeal airways and connectors				
			Sterilization of medical devices -				
			Microbiological methods Part_1:				
			Determination of a population of				
ISO 11737-1	2006-04	Ν	microorganisms on products	Y	N		
			Sterilization of medical devices				
			Microbiological methods Part_1:				
			Determination of a population of				
			microorganisms on products; Technical				
ISO 11737-1 Technic	al Corri 2007-05	Ν	Corrigendum_1	Y	N		
			Sterilization of medical devices	-			
			Microbiological methods Part_2: Tests				
			of sterility performed in the definition,				
			validation and maintenance of a				
ISO 11737-2	2009-11	Ν	sterilization process	Y	N		
				-			
			Lasers and laser-related equipment				
			Test method and classification for the				
			laser resistance of surgical drapes and/or				
			patient protective covers Part_1:				
ISO 11810-1	2005-02	Ν	Primary ignition and penetration				
	2000 02		Lasers and laser-related equipment				
			Test method and classification for the				
			laser-resistance of surgical drapes and/or				
			patient-protective covers Part_2:				
ISO 11810-2	2007-05	Ν	Secondary ignition				
	2007 00						1
			Acoustics Determination of sound immission				
			from sound sources placed close for the ear				
			Part_1: Technique using a microphone in a				
ISO 11904-1	2002-10	N	real ear (MIRE technique)				

			Urine-absorbing aids Part_1: Whole-product				1
ISO 11948-1	1996-11	Ν	testing				
130 11948-1	1990-11	IN	Dentistry Implants Clinical performance of				
ISO 11953	2010-06	Ν	hand torque instruments				
	2010 00						
			Ophthalmic optics Contact lenses and				
			contact lens care products Information				
ISO 11978	2000-03	N	supplied by the manufacturer				
130 11978	2000-03	IN	Ophthalmic implants Intraocular				
ISO 11979-1	2006-07	Ν	lenses Part_1: Vocabulary				
130 11979-1	2000-07	IN	Ophthalmic implants Intraocular				
ISO 11979-10	2006-08	Ν	lenses Part_10: Phakic intraocular lenses				
150 11979-10	2006-08	IN					
			Ophthalmic implants Intraocular				
100 44070 0	1000 10	N	lenses Part_2: Optical properties and				
ISO 11979-2	1999-12	N	test methods				
			Ophthalmic implants Intraocular				
			lenses Part_2: Optical properties and				
ISO 11979-2 Technical C	orri 2003-11	N	test methods; Technical Corrigendum_1				
			Ophthalmic implants Intraocular				
			lenses Part_3: Mechanical properties				
ISO 11979-3	2006-05	N	and test methods			 	
			Ophthalmic implants Intraocular				
			lenses Part_4: Labelling and				
ISO 11979-4	2008-12	N	information				
			Ophthalmic implants Intraocular				
ISO 11979-5	2006-06	N	lenses Part_5: Biocompatibility				
			Ophthalmic implants Intraocular				
			lenses Part_6: Shelf-life and transport				
ISO 11979-6	2007-07	N	stability				
			Ophthalmic implants Intraocular				
ISO 11979-7	2006-05	N	lenses Part_7: Clinical investigations	Y	N		
			Ophthalmic implants Intraocular				
			lenses Part_7: Clinical investigations;				
ISO 11979-7 AMD 1	2012-01	N	Amendment_1			 	
			Ophthalmic implants Intraocular				
			lenses Part_8: Fundamental				
ISO 11979-8	2006-07	N	requirements			 	
			Ophthalmic implants Intraocular				
			lenses Part_8: Fundamental				
ISO 11979-8 AMD 1	2011-05	N	requirements; Amendment_1			 	
			Ophthalmic implants Intraocular				
			lenses Part_9: Multifocal intraocular				
ISO 11979-9	2006-09	N	lenses				

			Ophthalmic optics Contact lenses and		
			contact lens care products Guidance		
ISO 11980	2009-10	Ν	for clinical investigations		
130 11900	2009-10	IN	Ophthalmic optics Contact lenses and		
			contact lens care products		
			Determination of physical compatibility of		
100 44004	0000 07		contact lens care products with contact		
ISO 11981	2009-07	N	lenses	 	
			Ophthalmic optics Contact lenses		
100 44005	4007.40	N	Ageing by exposure to UV and visible		
ISO 11985	1997-12	N	radiation (in vitro method)		
			Ophthalmic optics Contact lenses and		
			contact lens care products		
10.0 / / 00.0			Determination of preservative uptake and		
ISO 11986	2010-11	N	release		
			Ophthalmic optics Contact lenses		
ISO 11987	1997-12	N	Determination of shelf-life		
			Ophthalmic optics Contact lenses		
			Determination of shelf-life; Technical		
ISO 11987 Technical Co	orrige 1998-04	N	Corrigendum_1		
			Lasers and laser-related equipment		
			Determination of laser resistance of		
			tracheal tubes Part_1: Tracheal tube		
ISO 11990-1	2011-08	N	shaft		
			Lasers and laser-related equipment		
			Determination of laser resistance of		
			tracheal tubes Part_2: Tracheal tube		
ISO 11990-2	2010-07	N	cuffs		
			Health informatics Digital imaging and communication in medicine (DICOM) including		
ISO 12052	2006-11	Ν	workflow and data management		
130 12032	2000-11	IN	Acoustics Procedures for the measurement		
			of real-ear acoustical characteristics of hearing		
ISO 12124	2001-03	Ν	aids		
			Implants for surgery Mechanical testing		
			of implantable spinal devices Fatigue		
			test method for spinal implant assemblies		
ISO 12189	2008-05	Ν	using an anterior support		
	1		Medical gloves made from natural rubber	ľ	
			latex Determination of water-		
			extractable protein using the modified		
ISO 12243	2003-10	Ν	Lowry method		
			Tissue paper and tissue products Part_1:		1
ISO 12625-1	2011-08	Ν	General guidance on terms		
			Tissue paper and tissue products Part_12:		
100 40005 40	0040.04		Determination of tensile strength of perforated		
ISO 12625-12	2010-01	N	lines Calculation of perforation efficiency		

[			Tissue paper and tissue products - Part 3:			
			Determination of thickness, bulking thickness			
100 10005 0	2005 04	N				
ISO 12625-3	2005-04	N	and apparent bulk density Tissue paper and tissue products Part_4:			
100 40005 4	0005.04	N	Determination of tensile strength, stretch at			
ISO 12625-4	2005-04	N	break and tensile energy absorption		 	
100 40005 5	0005.04		Tissue paper and tissue products Part_5:			
ISO 12625-5	2005-04	N	Determination of wet tensile strength Tissue paper and tissue products - Part 6:			
190 40005 0	0005.00	N				
ISO 12625-6	2005-02	N	Determination of grammage			
100 40005 7	0007.00	N	Tissue paper and tissue products Part_7:			
ISO 12625-7	2007-03	N	Determination of optical properties		 	
			Tissue paper and tissue products Part_8:			
100 10005 0	0040.40		Water-absorption time and water-absorption			
ISO 12625-8	2010-12	N	capacity, basket-immersion test method		 	
100 10005 0	0005.05		Tissue paper and tissue products Part_9:			
ISO 12625-9	2005-05	N	Determination of ball burst strength		 	
			Ophthalmic optics Contact lenses			
ISO 12864	1997-12	N	Determination of scattered light			
ISO 12865	2006-07	N	Ophthalmic instruments Retinoscopes			
ISO 12866	1999-06	Ν	Ophthalmic instruments - Perimeters			
			Ophthalmic instruments Perimeters;			
ISO 12866 AMD 1	2008-11	Ν	Amendment 1			
ISO 12867	2010-06	N	Ophthalmic instruments Trial frames		 	
150 12867	2010-06	IN	Ophthalmic instruments Thai frames			
			Ophthalmic optics Spectacle frames			
ISO 12870	2004-08	N	Requirements and test methods			
			Implants for surgery Retrieval and			
			analysis of surgical implants - Part 1:			
ISO 12891-1	2011-05	Ν	Retrieval and handling			
			Retrieval and analysis of surgical			
			implants - Part 2: Analysis of retrieved			
ISO 12891-2	2000-02	N	· ,			
130 12091-2	2000-02	IN	metallic surgical implants			
			Retrieval and analysis of surgical			
			implants Part_3: Analysis of retrieved			
ISO 12891-3	2000-02	N	polymeric surgical implants			
			Retrieval and analysis of surgical			
			implants Part_4: Analysis of retrieved			
ISO 12891-4	2000-02	Ν	ceramic surgical implants			
			Health informatics Service architecture			
ISO 12967-1	2009-08	Ν	Part_1: Enterprise viewpoint			
	2000 00		Health informatics Service architecture			
ISO 12967-2	2009-08	Ν	Part_2: Information viewpoint			
			Health informatics - Service architecture -			
ISO 12967-3	2009-08	Ν	Part_3: Computational viewpoint			
	2000 00		Ophthalmic optics Contact lens care			
			products - Guidelines for determination			
100 10010	0044.05					
ISO 13212	2011-05	N	of shelf-life			

ISO 13294	1997-05	N	Dental handpieces Dental air-motors				
ISO 13295	2007-07	N	Dentistry Mandrels for rotary instruments				
			Implants for surgery Ceramic materials				
			based on yttria-stabilized tetragonal				
ISO 13356	2008-06	Ν	zirconia (Y-TZP)				
	2000 00		Periodontal curettes, dental scalers and				
ISO 13397-1	1995-12	Ν	excavators Part_1: General requirements				
			Dentistry Periodontal curettes, dental				
			scalers and excavators Part_2: Periodontal				
ISO 13397-2	2005-06	N	curettes of Gr-type				
100 40007 0	1000.00		Periodontal curettes, dental scalers and				
ISO 13397-3	1996-09	N	excavators Part_3: Dental scalers H-type Periodontal curettes, dental scalers and				
			excavators - Part 4: Dental excavators -				
ISO 13397-4	1997-12	Ν	Discoid type				
	1007 12						
			Surgical and dental hand instruments				
			Determination of resistance against				
ISO 13402	1995-08	N	autoclaving, corrosion and thermal exposure				
			Prosthetics and orthotics Categorization and				
100 1010			description of external orthoses and orthotic				
ISO 13404	2007-07	N	components			 	
			Prosthetics and orthostics Classification and				
			description of prosthetic components Part_1:				
ISO 13405-1	1996-10	Ν	Classification of prosthetic components				
	1000 10		Prosthetics and orthostics - Classification and				
			description of prosthetic components Part_2:				
			Description of lower-limb prosthetic				
ISO 13405-2	1996-10	N	components				
			Prosthetics and orthostics Classification and				
			description of prosthetic components Part_3:				
ISO 13405-3	1996-10	Ν	Description of upper-limb prosthetic components				
130 13405-3	1990-10	IN	components				
			Aseptic processing of health care				
ISO 13408-1	2008-06	Ν	products Part_1: General requirements	Y	N		
130 13400-1	2008-00	IN	Aseptic processing of health care	I	IN		
ISO 13408-2	2003-03	Ν	products Part_2: Filtration	Y	N		
130 13400-2	2003-03	IN	Aseptic processing of health care	I	IN		
ISO 13408-3	2006-09	Ν	products Part_3: Lyophilization	Y	N		
130 13400-3	2000-09	IN	Aseptic processing of health care	Ť	IN		+
			products Part_4: Clean-in-place				
100 10100 1	2005 11	N		×.	N		
ISO 13408-4	2005-11	N	technologies	Y	N		
			Acomtic processing of boolth acro				
180 12408 5	2006 44	NI	Aseptic processing of health care	V	N		
ISO 13408-5	2006-11	N	products Part_5: Sterilization in place	Y	N		

	<u>г</u>		Acceptic processing of bactth core				
180 13108 6	2005.00	N	Aseptic processing of health care	Y	N		
ISO 13408-6	2005-06	N	products Part_6: Isolator systems	Ý	N		
			Medical devices Quality management				
			systems Requirements for regulatory				
ISO 13485	2003-07	N	purposes	Y	N	AS ISO 13485	
			Medical devices Quality management				
			systems Requirements for regulatory				
ISO 13485 Technical Corrig	e2009-08	Ν	purposes; Technical Corrigendum_1	Y	Ν		
			Health informatics Electronic health record				
ISO 13606-1	2008-02	N	communication Part_1: Reference model				
			Health informatics Electronic health record				
100 10000 0	0000.40		communication Part_2: Archetype				
ISO 13606-2	2008-12	N	interchange specification Health informatics Electronic health record				
			communication Part_3: Reference				
ISO 13606-3	2009-02	Ν	archetypes and term lists				
130 13000-3	2009-02	IN	Health informatics Electronic health record				
			communication Part_5: Interface				
ISO 13606-5	2010-03	Ν	specification				
	2010 00		Ophthalmic optics Spectacle lenses				
ISO 13666	1998-08	Ν	Vocabulary				
	1000 00	11	Dentistry Reversible-irreversible hydrocolloid				
ISO 13716	1999-05	Ν	impression material systems				
	1000 00		Implants for surgery Hydroxyapatite				
ISO 13779-1	2008-10	Ν	Part_1: Ceramic hydroxyapatite				
	2000 10		Implants for surgery Hydroxyapatite				
ISO 13779-2	2008-10	Ν	Part_2: Coatings of hydroxyapatite				
100 13773-2	2000-10	IN	Implants for surgery Hydroxyapatite				
			Part_3: Chemical analysis and				
100 40770 0	0000.00	NI	characterization of crystallinity and phase				
ISO 13779-3	2008-02	N	purity				
			Implants for surgery Hydroxyapatite				
			Part_4: Determination of coating				
ISO 13779-4	2002-05	N	adhesion strength				
			Poly(L-lactide) resins and fabricated				
			forms for surgical implants In vitro				
ISO 13781	1997-02	N	degradation testing				
			Implants for surgery Metallic materials				
			Unalloyed tantalum for surgical implant				
ISO 13782	1996-12	Ν	applications				
ISO 13897	2003-02	Ν	Dentistry Amalgam capsules				
			Dentistry Amalgam capsules; Technical				
ISO 13897 Technical Corrigence	1 2003-12	Ν	Corrigendum_1				
			Pen systems Part_1: Glass cylinders				
ISO 13926-1	2004-11	Ν	for pen-injectors for medical use				

			Pen systems Part_2: Plunger stoppers				
ISO 13926-2	2011-04	Ν	for pen-injectors for medical use				
100 10020 2	2011.04		Concentrates for haemodialysis and				
ISO 13958	2009-04	Ν	related therapies				
			Water for haemodialysis and related				
ISO 13959	2009-04	Ν	therapies				
			Cardiovascular implants and				
ISO 13960	2010-07	Ν	extracorporeal systems Plasmafilters				
			Clinical investigation of medical devices				
			for human subjects Good clinical				
ISO 14155	2011-02	N	practice	Y	N	AS ISO 14155-1	
			Clinical investigation of medical devices				
			for human subjects Good clinical				
ISO 14155 Technical	Corrige 2011-07	N	practice; Technical Corrigendum_1				
			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				
ISO 14160	2011-07	Ν	control of a sterilization process for medical devices	Y	N	AS ISO 14160	
150 14160	2011-07	IN	Sterilization of health care products	ř	IN	AS 150 14160	
			Biological indicators Guidance for the				
			selection, use and interpretation of				
ISO 14161	2009-09	Ν	results				
ISO 14233	2003-03	N	Dentistry Polymer-based die materials				
			Implants for surgery Wear of total hip-				
			joint prostheses Part_1: Loading and				
			displacement parameters for wear-testing				
			machines and corresponding				
ISO 14242-1	2012-01	Ν	environmental conditions for test				
			Implants for surgery Wear of total hip				
			joint prostheses Part_2: Methods of				
ISO 14242-2	2000-09	N	measurement				
			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				
ISO 14242-3	2009-03	N	for test				

					r	r
			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			
ISO 14243-1	2009-11	N	for test			
			Implants for surgery Wear of total knee-			
			joint prostheses Part_2: Methods of			
ISO 14243-2	2009-11	N	measurement			
l						
			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			
ISO 14243-3	2004-09	N	for test			
			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			
ISO 14243-3 Technica	al Corri 2006-02	Ν	for test			
ISO 14356	2003-03	N	Dentistry Duplicating material			
			Tracheal tubes designed for laser			
			surgery Requirements for marking and			
ISO 14408	2005-06	N	accompanying information			
			Ophthalmic optics Contact lenses and			
			contact lens care products			
ISO 14534	2011-04	Ν	Fundamental requirements			
			Non-active surgical implants Implants			
			for osteosynthesis Particular			
ISO 14602	2010-04	Ν	requirements			
			Non-active surgical implants Mammary			
ISO 14607	2007-02	Ν	implants Particular requirements			
			Non-active surgical implants General			
ISO 14630	2008-01	Ν	requirements			
			Implants for surgery Active implantable			
			medical devices Part_1: General			
			requirements for safety, marking and for			
			information to be provided by the			
ISO 14708-1	2000-11	Ν	manufacturer			
			Implants for surgery Active implantable			
			medical devices Part_2: Cardiac			
ISO 14708-2	2005-10	Ν	pacemakers			

			Implente for europris Active implementable				
			Implants for surgery Active implantable				
100 4 4700 0	0000 44	N	medical devices Part_3: Implantable				
ISO 14708-3	2008-11	N	neurostimulators				
			Implants for surgery Active implantable				
100 4 4700 4	0000 44		medical devices Part_4: Implantable				
ISO 14708-4	2008-11	N	infusion pumps				
			Implants for surgery Active implantable				
			medical devices Part_5: Circulatory				
ISO 14708-5	2010-02	N	support devices				
			Implants for surgery Active implantable				
			medical devices Part_6: Particular				
			requirements for active implantable				
			medical devices intended to treat				
			tachyarrhythmia (including implantable				
ISO 14708-6	2010-03	N	defibrillators)				
			Ophthalmic optics Contact lens care				
			products Microbiological requirements				
			and test methods for products and				
			regimens for hygienic management of				
ISO 14729	2001-04	N	contact lenses				
			Ophthalmic optics Contact lens care				
			products Microbiological requirements				
			and test methods for products and				
			regimens for hygienic management of				
ISO 14729 AMD 1	2010-10	N	contact lenses; Amendment_1				
			Ophthalmic optics Contact lens care				
			products Antimicrobial preservative				
			efficacy testing and guidance on				
ISO 14730	2000-09	N	determining discard date				
100 / 100 /			Dentistry Implants Dynamic fatigue test for				
ISO 14801	2007-11	N	endosseous dental implants				
			<b>–</b> – – – – – – – – – – – – – – – – – –				
			Implants for surgery Total knee-joint				
			prostheses Part_1: Determination of				
ISO 14879-1	2000-06	N	endurance properties of knee tibial trays				
			Ophthalmic optics Spectacle lenses				
			Fundamental requirements for uncut				
ISO 14889	2003-05	N	finished lenses				
			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				
ISO 14937	2009-10	N	medical devices	Y	Ν		
			Implants for surgery Two-part addition-				
ISO 14949	2001-10	N	cure silicone elastomers				

			Medical devices Application of risk				
ISO 14971	2007-03	Ν	management to medical devices	Y	N		
100 1437 1	2007-03	IN	Indiagement to medical devices	1			
			Sterile obturators for single use with over-				
ISO 14972	1998-12	Ν	needle peripheral intravascular catheters				
130 14972	1990-12	IN	Anaesthetic and respiratory equipment				
100 45004	0040.00	NI					
ISO 15001	2010-06	N	Compatibility with oxygen				
			Flow-metering devices for connection to				
100 15000			terminal units of medical gas pipeline				
ISO 15002	2008-07	N	systems				
			Ophthalmic instruments Fundamental				
			requirements and test methods Part_1:				
			General requirements applicable to all				
ISO 15004-1	2006-06	N	ophthalmic instruments				
			Ophthalmic instruments Fundamental				
			requirements and test methods Part_2:				
ISO 15004-2	2007-02	N	Light hazard protection				
			Disposable hanging devices for				
			transfusion and infusion bottles				
ISO 15010	1998-06	N	Requirements and test methods				
			Prostheses Structural testing of hip				
ISO 15032	2000-04	Ν	units				
			Dental elevators Part_1: General				
ISO 15087-1	1999-11	N	requirements				
			Dental elevators Part_2: Warwick James				
ISO 15087-2	2000-04	N	elevators				
ISO 15087-3	2000-05	N	Dental elevators Part_3: Cryer elevators				
ISO 15087-4	2000-05	Ν	Dental elevators Part_4: Coupland elevators				
ISO 15087-5	2000-05	N	Dental elevators Part_5: Bein elevators				
ISO 15087-5	2000-05	N	Dental elevators Part_6: Flohr elevators				
150 15087-6	2000-05	IN	Dental tweezers Part_1: General				
ISO 15098-1	1999-10	Ν	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				
100 10000-0	2000-02	11	Self-adhesive hanging devices for				
			infusion bottles and injection vials -				
ISO 15137	2005-07	Ν	Requirements and test methods				
100 10101	2000-07	IN	Implants for surgery Metal				
100 45440 4	2002.00	<b>K</b> 1	intramedullary nailing systems Part_1:				
ISO 15142-1	2003-08	N	Intramedullary nails			 	
			Implants for surgery Metal				
100 45440 0	0000.00		intramedullary nailing systems Part_2:				
ISO 15142-2	2003-08	N	Locking components				

			Implants for surgery Metal			
			intramedullary nailing systems Part_3:			
			Connection devices and reamer diameter			
ISO 15142-3	2003-08	Ν	measurements			
	2000 00		In vitro diagnostic medical devices			
			Measurement of quantities in samples of			
			biological origin Requirements for			
			content and presentation of reference			
ISO 15193	2009-05	Ν	measurement procedures			
			In vitro diagnostic medical devices			
			Measurement of quantities in samples of			
			biological origin Requirements for			
			certified reference materials and the			
ISO 15194	2009-05	Ν	content of supporting documentation			
			In vitro diagnostic test systems			
			Requirements for blood-glucose monitoring			
			systems for self-testing in managing diabetes			
ISO 15197	2003-05	N	mellitus			
			Clinical laboratory medicine In vitro			
			diagnostic medical devices Validation			
100 45400	0004.07	N	of user quality control procedures by the			
ISO 15198	2004-07	N	manufacturer Medical devices Symbols to be used			 
			with medical device labels, labelling and			
			information to be supplied Part_1:			
ISO 15223-1	2007-04	N	General requirements			
150 15225-1	2007-04	IN	General requirements			
			Medical devices Symbols to be used			
			with medical device labels, labelling and			
			information to be supplied Part_1:			
ISO 15223-1 AMD 1	2008-06	Ν	General requirements; Amendment_1			
			Medical devices Symbols to be used			
			with medical device labels, labelling, and			
			information to be supplied Part_2:			
			Symbol development, selection and			
ISO 15223-2	2010-01	Ν	validation			
			Medical devices Quality management			
			Medical device nomenclature data			
ISO 15225	2010-05	Ν	structure			
			Ophthalmic optics and instruments Optical			
ISO 15253	2000-09	N	devices for enhancing low vision			 
			Ophthalmic optics and instruments			
			Electro-optical devices for enhancing low			
ISO 15254	2009-07	N	vision			
			Implants for surgery Requirements for			
ISO 15374	1998-08	N	production of forgings			

			Medical infusion bottles Suspension			T
100 45075	2010-06	NI	devices for multiple use Requirements and test methods			
ISO 15375	2010-06	N	and test methods			
			Primary packaging materials for medicinal			
			products Particular requirements for the			
			application of ISO_9001:2008, with reference			
ISO 15378	2011-11	Ν	to Good Manufacturing Practice_(GMP)			
			Dental handpieces Air-powered scalers and			
ISO 15606	1999-12	N	scaler tips			
100 45004	0011.00		Urine-absorbing aids General guidelines on			
ISO 15621	2011-02	N	evaluation			
ISO 1563	1990-09	N	Dental alginate impression material			
ISO 1564	1995-11	N	Dental aqueous impression materials based on agar			
130 1304	1990-11	IN	Ullagai			
			Cardiovascular implants and artificial			
			organs Hard-shell cardiotomy/venous			
			reservoir systems (with/without filter) and			
ISO 15674	2009-04	Ν	soft venous reservoir bags			
130 13074	2003-04	IN	Cardiovascular implants and artificial			
			organs Cardiopulmonary bypass			
ISO 15675	2009-04	Ν	systems Arterial blood line filters			
150 15075	2003-04		Cardiovascular implants and artificial			
			organs Requirements for single-use			
			tubing packs for cardiopulmonary bypass			
			and extracorporeal membrane			
ISO 15676	2005-07	Ν	oxygenation (ECMO)			
100 10070	2000 07		Plastic containers for intravenous			
ISO 15747	2010-04	Ν	injections			
100 10141	2010 04		Ophthalmic instruments			
			Endoilluminators Fundamental			
			requirements and test methods for optical			
ISO 15752	2010-01	Ν	radiation safety			
	2010 01	••	Medical infusion equipment Plastics			
			caps with inserted elastomeric liner for			
			containers manufactured by the blow-fill-			
ISO 15759	2005-04	Ν	seal (BFS) process			
			Ophthalmic implants Ophthalmic			
ISO 15798	2010-01	Ν	viscosurgical devices			
-			Implants for surgery Copolymers and			
			blends based on polylactide In vitro			
ISO 15814	1999-11	Ν	degradation testing			
ISO 15841	2006-10	N	Dentistry Wires for use in orthodontics			
ISO 15854	2005-07	N	Dentistry Casting and baseplate waxes			

	T		Sterilization of health care products	I	1	1	
			Chemical indicators - Guidance for				
			selection, use and interpretation of				
ISO 15882	2008-09	Ν	results				
130 13662	2008-09	IN	Washer-disinfectors Part_1: General				
			requirements, terms and definitions and				
ISO 15883-1	2006-04	Ν	tests				
130 13663-1	2000-04	IN	Washer-disinfectors Part_2:				
l			Requirements and tests for washer-				
			disinfectors employing thermal				
			disinfection for surgical instruments,				
			anaesthetic equipment, bowls, dishes,				
ISO 15883-2	2006-04	Ν	receivers, utensils, glassware, etc.				
	2000 01	••					
			Washer-disinfectors Part_3:				
			Requirements and tests for washer-				
			disinfectors employing thermal				
ISO 15883-3	2006-04	Ν	disinfection for human waste containers				
			Washer-disinfectors Part_4:				
			Requirements and tests for washer-				
			disinfectors employing chemical				
ISO 15883-4	2008-05	N	disinfection for thermolabile endoscopes				
			Washer-disinfectors Part_6:				
			Requirements and tests for washer-				
			disinfectors employing thermal				
			disinfection for non-invasive, non-critical				
			medical devices and healthcare				
ISO 15883-6	2011-04	N	equipment Dentistry Casting investments and refractory				
ISO 15912	2006-10	N	die materials				
			Dentistry Casting investments and refractory				
			die materials; Amendment_1: Requirement				
	0011.07		and test method for adequacy of expansion of				
ISO 15912 AMD 1	2011-07	N	Type_1 and Type_2 materials Urine-absorbing aids Basic principles for				
			evaluation of single-use adult-incontinence-				
			absorbing aids from the perspective of users				
ISO 16021	2000-11	Ν	and caregivers				
			Ophthalmic optics Specifications for				
			single-vision ready-to-wear near-vision				
ISO 16034	2002-02	N	spectacles				
			Ophthalmic optics Specifications for				
			single-vision ready-to-wear near- vision				
ISO 16034 Technical C	Corrige 2006-08	N	spectacles; Technical Corrigendum_1				

r			Rubber condoms for clinical trials -	Γ		ſ
ISO 16037	2002-05	Ν	Measurement of physical properties			
130 10037	2002-03	IN	Rubber condoms for clinical trials			
			Measurement of physical properties;			
ISO 16037 AMD 1	2011-02	Ν	Amendment 1			
130 10037 AMD 1	2011-02	IN	Rubber condoms Guidance on the use of			
			ISO_4074 in the quality management of			
ISO 16038	2005-11	Ν	natural rubber latex condoms			
130 10038	2003-11	IN	Implants for surgery Minimum data sets			
100 10051	2000 42	N				
ISO 16054	2000-12	N	for surgical implants Dentistry Required elements for codification	 		
100 40050	0007.00	N				
ISO 16059	2007-08	N	used in data exchange	 		
			Instrumentation for use in association			
			with non-active surgical implants			
ISO 16061	2008-12	N	General requirements			
			Technical aids for persons with disability			
			Environmental control systems for daily			
ISO 16201	2006-10	Ν	living			
	2000.0		Ophthalmic optics Information			
			interchange for ophthalmic optical			
100 10081	2000 02	N				
ISO 16284	2006-03	N	equipment	 		
			Aids for ostomy and incontinence Irrigation			
100 10201	2002 40	Ν	sets Requirements and test methods			
ISO 16391	2002-10	IN		 		
			Implants for surgery Acrylic resin			
			cement Flexural fatigue testing of			
			acrylic resin cements used in			
ISO 16402	2008-05	N	orthopaedics			
			Dentistry Oral hygiene products Oral			
ISO 16408	2004-04	N	rinses			
			Dentistry Oral hygiene products Manual			
ISO 16409	2006-10	N	interdental brushes	 		
			Dentistry Oral hygiene products Manual			
ISO 16409 AMD 1	2010-02	N	interdental brushes; Amendment_1	 		
			Implants for surgery Test solutions and			
			environmental conditions for static and			
			dynamic corrosion tests on implantable			
ISO 16428	2005-04	Ν	materials and medical devices			
	2000 01		Implants for surgery Measurements of			
			open-circuit potential to assess corrosion		1	
			behaviour of metallic implantable		1	
			materials and medical devices over		1	
ISO 16429	2004-07	N	extended time periods			
			Tracheobronchial tubes Sizing and			
ISO 16628	2008-11	Ν	marking			
			Ophthalmic implants Irrigating solutions			
ISO 16671	2003-05	Ν	for ophthalmic surgery			
	2000 00				1	1

			Ophthalmic implants Ocular			
ISO 16672	2003-02	Ν	endotamponades			
100 10072	2000 02	11	Wheelchair seating Part_1: Vocabulary,		1	
			reference axis convention and measures for			
			body segments, posture and postural support			
ISO 16840-1	2006-03	Ν	surfaces			
			Wheelchair seating Part_2: Determination of			
			physical and mechanical characteristics of			
			devices intended to manage tissue integrity			
ISO 16840-2	2007-07	Ν	Seat cushions			
			Wheelchair seating Part_3: Determination of			
			static, impact and repetitive load strengths for			
ISO 16840-3	2006-07	N	postural support devices			
			Wheelchair seating Part_4: Seating systems			
ISO 16840-4	2009-03	N	for use in motor vehicles			
			Leolth information Dublic key infor-two-two-			
100 47000 4	0000.00		Health informatics Public key infrastructure			
ISO 17090-1	2008-02	N	Part_1: Overview of digital certificate services Health informatics Public key infrastructure			
ISO 17090-2	2008-02	Ν	Part_2: Certificate profile			
130 17090-2	2000-02	IN	Health informatics Public key infrastructure			
			Part_3: Policy management of certification			
ISO 17090-3	2008-02	Ν	authority			
100 11030 0	2000 02		Health informatics Vocabulary for			
ISO 17115	2007-07	Ν	terminological systems			
	2001 01		Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_1: Determination			
ISO 17190-1	2001-12	Ν	of_pH			
			Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_10: Determination			
			of extractable polymer content by			
ISO 17190-10	2001-12	N	potentiometric titration			
			Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
ISO 17190-11	2001-12	Ν	absorbent materials Part_11: Determination of content of respirable particles			
130 17 190-11	2001-12	IN			1	
			Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based		1	
			absorbent materials Part_2: Determination			
ISO 17190-2	2001-12	Ν	of amount of residual monomers			
			Urine absorbing aids for incontinence Test			
			methods for characterizing polymer-based		1	
			absorbent materials Part_3: Determination			
			of particle size distribution by sieve			
ISO 17190-3	2001-12	Ν	fractionation			

				[	-		
			Urine-absorbing aids for incontinence Test				
			methods for characterizing polymer-based				
			absorbent materials Part_4: Determination				
ISO 17190-4	2001-12	N	of moisture content by mass loss upon heating				
			Urine-absorbing aids for incontinence Test				
			methods for characterizing polymer-based				
			absorbent materials Part_5: Gravimetric				
100 17100 5	2001-12	N	determination of free swell capacity in saline				
ISO 17190-5	2001-12	N	solution				
l I			Urine-absorping aids for incontinence Test				
			methods for characterizing polymer-based				
			absorbent materials Part_6: Gravimetric				
			determination of fluid retention capacity in				
ISO 17190-6	2001-12	N	saline solution after centrifugation				
			Ultime choose in a side for incontinuous. The				
			Urine-absorbing aids for incontinence Test methods for characterizing polymer-based				
			absorbent materials Part_7: Gravimetric				
ISO 17190-7	2001-12	N	determination of absorption under pressure				
100 17 190-7	2001-12	IN	Urine-absorping aids for incontinence - Test				
			methods for characterizing polymer-based				
			absorbent materials Part_8: Gravimetric				
ISO 17190-8	2001-12	Ν	determination of flowrate				
			Urine-absorbing aids for incontinence Test				
			methods for characterizing polymer-based				
			absorbent materials Part_9: Gravimetric				
ISO 17190-9	2001-12	N	determination of density				
			Urine-absorbing aids for incontinence Test				
			methods for characterizing polymer-based absorbent materials Part_9: Gravimetric				
			determination of density; Technical				
ISO 17190-9 Technical	Corrigen 2002-10	Ν	Corrigendum_1				
	0011g012002 10						
			Urine-absorbing aids for incontinence				
			Measurement of airborne respirable				
			polyacrylate superabsorbent materials				
			Determination of dust in collection cassettes				
ISO 17191	2004-02	N	by sodium atomic absorption spectrometry				
			Health informatics Messages and				
100 17400	2004 42	N	communication Web access to DICOM				
ISO 17432	2004-12	N	persistent objects				
			Sleep apnoea breathing therapy				
100 47540 4	0007.40	N	Part_1: Sleep apnoea breathing therapy				
ISO 17510-1	2007-10	N	equipment				
			Sleep apnoea breathing therapy				
100 17510 0	0007.40		Part_2: Masks and application				
ISO 17510-2	2007-10	N	accessories				

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			In vitro diagnostic medical devices				
			Measurement of quantities in biological				
			samples Metrological traceability of				
			values assigned to calibrators and control				
ISO 17511	2003-08	N	materials				
			Clinical laboratory testing and in vitro				
l			medical devices Requirements for in				
l			vitro monitoring systems for self-testing of				
ISO 17593	2007-04	Ν	oral anticoagulant therapy				
			Sterilization of medical devices -				
			Information to be provided by the				
			manufacturer for the processing of				
ISO 17664	2004-03	Ν	resterilizable medical devices	Y	N		
100 17004	2004-03		Sterilization of health care products	I			
			Moist heat Part_1: Requirements for				
			the development, validation and routine				
100 17005 1			control of a sterilization process for				
ISO 17665-1	2006-08	N	medical devices	Y	N		
			Wear of implant materials Polymer and				
			metal wear particles Isolation and				
ISO 17853	2011-03	N	characterization				
			Dentistry Shanks for rotary instruments				
ISO 1797-1	2011-08	N	Part_1: Shanks made of metals				
100 1707 0	(000.00		Dental rotary instruments; shanks; part_2:				
ISO 1797-2	1992-02	N	shanks made of plastics				-
ISO 18084	2011-09	N	Press tools for tablets Punches and dies				
100 40404	0000.40	N	Health informatics Integration of a reference				
ISO 18104	2003-12	N	terminology model for nursing				
			In vitro diagnostic medical devices				
			Information supplied by the manufacturer				
			(labelling) Part_1: Terms, definitions				
ISO 18113-1	2009-12	N	and general requirements				
			In vitro diagnostic medical devices				
			Information supplied by the manufacturer				
			(labelling) Part_2: In vitro diagnostic				
ISO 18113-2	2009-12	Ν	reagents for professional use				
			In vitro diagnostic medical devices				
			Information supplied by the manufacturer				
			(labelling) Part_3: In vitro diagnostic				
ISO 18113-3	2009-12	Ν	instruments for professional use				

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			
			In vitro diagnostic medical devices			
			Information supplied by the manufacturer			
			(labelling) Part_5: In vitro diagnostic			
ISO 18113-5	2009-12	N	instruments for self-testing			
			In vitro diagnostic medical devices			
			Measurement of quantities in biological			
			samples Metrological traceability of			
			values for catalytic concentration of			
			enzymes assigned calibrators and control			
ISO 18153	2003-08	N	materials			
			Implants for surgery Wear of total			
			intervertebral spinal disc prostheses			
			Part_1: Loading and displacement			
			parameters for wear testing and			
100 40400 4	0011.00		corresponding environmental conditions			
ISO 18192-1	2011-03	N	for test			
			Implants for surgery Wear of total intervertebral spinal disc prostheses			
ISO 18192-2	2010-06	Ν	Part_2: Nucleus replacements			
130 10192-2	2010-00	IN	Health Informatics Messages and			
			communication Format of length limited			
ISO 18232	2006-04	Ν	globally unique string identifiers			
			Health informatics Requirements for an			
ISO 18308	2011-04	N	electronic health record architecture			
			Ophthalmic optics Contact lenses			
			Part_1: Vocabulary, classification system			
100 10000 1			and recommendations for labelling			
ISO 18369-1	2006-08	Ν	specifications			
			Ophthalmic optics Contact lenses			
			Part_1: Vocabulary, classification system and recommendations for labelling			
	2000.02	N	0			
ISO 18369-1 AMD 1	2009-02	Ν	specifications; Amendment_1			
ISO 18369-2	2006-08	Ν	Ophthalmic optics Contact lenses Part_2: Tolerances			
100 10009-2	2000-00	ſN	Ophthalmic optics Contact lenses			
ISO 18369-3	2006-08	Ν	Part_3: Measurement methods			
100 10009-0	2000-00	IN	Ophthalmic optics Contact lenses			
			Part_4: Physicochemical properties of			
ISO 18369-4	2006-08	N	contact lens materials			

			Sterilization of health care products				
			Biological and chemical indicators Test				
ISO 18472	2006-06	Ν	equipment				
130 18472	2000-00	IN	equipment				
			Transportable liquid ovugan ovatame for				
100 40777	0005.00		Transportable liquid oxygen systems for				
ISO 18777	2005-02	N	medical use Particular requirements	-	_	-	
			Respiratory equipment Infant monitors				
ISO 18778	2005-02	N	Particular requirements				
			Medical devices for conserving oxygen				
			and oxygen mixtures Particular				
ISO 18779	2005-02	N	requirements				
			Health informatics Clinical analyser				
			interfaces to laboratory information systems				
ISO 18812	2003-03	N	Use profiles				
			In vitro diagnostic medical devices				
			Information supplied by the manufacturer				
			with in vitro diagnostic reagents for				
ISO 19001	2002-11	Ν	staining in biology				
			Rail systems for supporting medical				
ISO 19054	2005-07	Ν	equipment				
ISO 1942	2009-12	Ν	Dentistry Vocabulary				
			Ophthalmic instruments - Corneal				
ISO 19980	2005-08	Ν	topographers				
			Aerosol drug delivery device design				
			verification Requirements and test				
ISO 20072	2009-08	Ν	methods				
	2000 00		Dentistry Manual toothbrushes General				
ISO 20126	2012-01	Ν	requirements and test methods				
			Dentistry Powered toothbrushes General				
ISO 20127	2005-03	Ν	requirements and test methods				
			Implants for surgery Metallic materials				
			Classification of microstructures for				
ISO 20160	2006-05	Ν	alpha+beta titanium alloy bars				
			Health informatics Health cards General				
ISO 20301	2006-11	Ν	characteristics				
			Health informatics Health cards				
			Numbering system and registration procedure				
ISO 20302	2006-12	Ν	for issuer identifiers				

	1					
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			
			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			
ISO 20776-2	2007-07	N	antimicrobial susceptibility test devices			
			Dentistry Base polymers Part_1: Denture			
ISO 20795-1	2008-08	Ν	base polymers			
			Dentistry Base polymers Part_1: Denture			
ISO 20795-1 Technical Corrige	n 2009-02	N	base polymers; Technical Corrigendum_1 Dentistry - Base polymers - Part 2:		 	
ISO 20795-2	2010-03	N	Orthodontic base polymers			
	2010 00		Sterilization of health care products Dry			
			heat Requirements for the			
			development, validation and routine			
			control of a sterilization process for			
ISO 20857	2010-08	N	medical devices			
			Health informatics Harmonized data types			
ISO 21090	2011-02	N	for information interchange			
			Medical gloves Determination of			
ISO 21171	2006-05	N	removable surface powder			
			Dentistry Materials used for dental equipment surfaces - Determination of			
ISO 21530	2004-06	N	resistance to chemical disinfectants			
100 21000	2004-00	IN IN	Dentistry Graphical symbols for dental			
ISO 21531	2009-02	N	instruments			
			Dentistry Reusable cartridge syringes			
ISO 21533	2003-06	N	intended for intraligamentary injections			
			Dentistry Reusable cartridge syringes			
ISO 21533 Technical Corrigence	1 2000 12	N	intended for intraligamentary injections; Technical Corrigendum_1			
150 21533 Technical Corrigenc	12009-12	IN				
			Non-active surgical implants Joint replacement implants Particular			
ISO 21534	2007-10	N	requirements			
130 21334	2007-10	IN				

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

			Madiaal daviana utilizing animal tiaguag			
			Medical devices utilizing animal tissues			
			and their derivatives Part_1:			
ISO 22442-1	2007-12	N	Application of risk management			
			Medical devices utilizing animal tissues			
			and their derivatives Part_2: Controls			
ISO 22442-2	2007-12	N	on sourcing, collection and handling			
			Medical devices utilizing animal tissues			
			and their derivatives Part_3: Validation			
			of the elimination and/or inactivation of			
			viruses and transmissible spongiform			
ISO 22442-3	2007-12	N	encephalopathy (TSE) agents			
			External limb prostheses and external			
			orthoses Requirements and test			
ISO 22523	2006-10	N	methods			
			Clothing for protection against infectious			
			agents - Medical face masks - Test method			
			for resistance against penetration by synthetic			
SO 22609	2004-12	Ν	blood (fixed volume, horizontally projected)			
	2001.12		Surgical drapes, gowns and clean air			
			suits, used as medical devices, for			
			patients, clinical staff and equipment			
			Test method to determine the resistance			
ISO 22610	2006-07	Ν	to wet bacterial penetration			
100 22010	2000-01		Clothing for protection against infectious			
			agents Test method for resistance to			
ISO 22612	2005-03	Ν	dry microbial penetration			
100 22012	2003-03	IN	Dentistry - Metallic materials for fixed and			
ISO 22674	2006-11	Ν	removable restorations and appliances			
			Prosthetics Testing of ankle-foot			
			devices and foot units Requirements			
ISO 22675	2006-10	Ν	and test methods			
ISO 22715	2006-04	N	Cosmetics Packaging and labelling			
100 221 10	2000 01		Cosmetics Good Manufacturing Practices			
			(GMP) Guidelines on Good Manufacturing			
ISO 22716	2007-11	Ν	Practices			
			Dentistry Implantable materials for bone			
			filling and augmentation in oral and			
			maxillofacial surgery Contents of a technical			
ISO 22794	2007-07	N	file			
			Dentistry Membrane materials for guided			
100 00000	0004.00	N	tissue regeneration in oral and maxillofacial			
ISO 22803	2004-09	N	surgery Contents of a technical file Health informatics - Guidelines on data			
			protection to facilitate trans-border flows of			
ISO 22857	2004-04	N	personal health information			
00 22001	2004-04	IN	personal health information		l	Į

r			Implanta for ourgany In vitro avaluation		T		ſ
			Implants for surgery In vitro evaluation				
100 00047	0007.00	NI	for apatite-forming ability of implant				
ISO 23317	2007-06	N	materials				
			Draathing quaters filters for an aathatia				
			Breathing system filters for anaesthetic				
100 00000 /			and respiratory use Part_1: Salt test				
ISO 23328-1	2003-08	N	method to assess filtration performance				
			Breathing system filters for anaesthetic				
			and respiratory use Part_2: Non-				
ISO 23328-2	2002-10	N	filtration aspects				
			Male condoms Requirements and test				
ISO 23409	2011-02	Ν	methods for condoms made from synthetic materials				
150 23409	2011-02	IN					
			Guidance for the preparation and quality				
100 00500	0011.05		management of fluids for haemodialysis				
ISO 23500	2011-05	N	and related therapies Assistive products for blind and vision-				
			impaired persons Tactile walking surface				
ISO 23599	2012-03	N	indicators				
100 20099	2012-03	IN	Assistive products for persons with vision				
			impairments and persons with vision and				
			hearing impairments Acoustic and tactile				
ISO 23600	2007-11	Ν	signals for pedestrian traffic lights				
			In vitro diagnostic medical devices				
			Evaluation of stability of in vitro				
ISO 23640	2011-12	Ν	diagnostic reagents				
			Anaesthetic and respiratory equipment				
			Peak expiratory flow meters for the				
			assessment of pulmonary function in				
ISO 23747	2007-07	Ν	spontaneously breathing humans				
-							
			Sharps injury protection Requirements				
			and test methods Sharps protection				
			features for single-use hypodermic				
			needles, introducers for catheters and				
ISO 23908	2011-06	Ν	needles used for blood sampling				
	2011 00			1		1	
			Ophthalmic optics and instruments				
ISO 24157	2008-07	Ν	Reporting aberrations of the human eve				
ISO 24214	2006-07	N	Skin barrier for ostomy aids - Vocabulary				
100 272 17	2000-11	IN	Dentistry Mercury and alloys for dental				
ISO 24234	2004-10	Ν	amalgam				
	2001.00		Dentistry Mercury and alloys for dental				
			amalgam Amendment_1: Requirements for				
			marking and manufacturer's instructions				
ISO 24234 AMD 1	2011-08	Ν	concerning mercury				

			Tipe for againtive products for wellking	1		
			Tips for assistive products for walking			
			Requirements and test methods Part_1:			
ISO 24415-1	2009-04	N	Friction of tips			
			Tips for assistive products for walking			
			Requirements and test methods Part_2:			
ISO 24415-2	2011-08	N	Durability of tips for crutches			
			Ergonomics Accessible design Auditory			
ISO 24500	2010-10	N	signals for consumer products			
			Ergonomics Accessible design Sound			
			pressure levels of auditory signals for			
ISO 24501	2010-12	N	consumer products			
			Ergonomics Accessible design			
			Specification of age-related luminance			
ISO 24502	2010-12	N	contrast for coloured light			
			Ergonomics Accessible designTactile dots			
ISO 24503	2011-01	Ν	and bars on consumer products			
			Sterilization of medical devices Low			1
			temperature steam and formaldehyde			
			Requirements for development,			
			validation and routine control of a			
10.0 07.40.4						
ISO 25424	2009-09	N	sterilization process for medical devices			
			Cardiovascular implants Endovascular			
			devices Part_1: Endovascular			
ISO 25539-1	2003-03	N	prostheses			
			Cardiovascular implants Endovascular			
			devices Part_1: Endovascular			
			prostheses; Amendment 1: Test			
ISO 25539-1 AMD 1	2005-07	Ν	methods			
150 25539-1 AIVID 1	2005-07	IN	Internous			
			Cardiovascular implants Endovascular			
ISO 25539-2	2008-09	N	devices Part_2: Vascular stents			
						1
			Cardiovascular implants Endovascular			
ISO 25539-3	2011-12	Ν	devices - Part 3: Vena cava filters			
	-		Health informatics - Genomic Sequence			
ISO 25720	2009-08	Ν	Variation Markup Language (GSVML)			
			Female condoms Requirements and test			
ISO 25841	2011-07	Ν	methods			
			Water treatment equipment for			
			haemodialysis applications and related			1
ISO 26722	2009-04	N	therapies			
130 20/22	2009-04	N				
			Anaesthetic and respiratory equipment			1
			Spirometers intended for the			
			measurement of time forced expired			1
ISO 26782	2009-07	Ν	volumes in humans			

			Anaesthetic and respiratory equipment			
			Spirometers intended for the			
			measurement of time forced expired			
			volumes in humans; Technical			
ISO 26782 Technical	Corrige 2009-11	N	Corrigendum_1			
			Anaesthetic and respiratory equipment			
			User-applied labels for syringes			
			containing drugs used during			
			anaesthesia Colours, design and			
ISO 26825	2008-08	N	performance			
			Dentistry Brackets and tubes for use in			
ISO 27020	2010-12	N	orthodontics			
			Cardiac rhythm management devices			
			Symbols to be used with cardiac rhythm			
			management device labels, and			
			information to be supplied General			
ISO 27185	2012-02	N	requirements			
			Active implantable medical devices			
			Four-pole connector system for			
			implantable cardiac rhythm management			
			devices Dimensional and test			
ISO 27186	2010-03	Ν	requirements			
			Anaesthetic and respiratory equipment			
ISO 27427	2010-03	Ν	Nebulizing systems and components			
100 21 121	2010 00		Health informatics Information security			
ISO 27799	2008-07	Ν	management in health using ISO/IEC_2702			
ISO 28158	2010-07	N	Dentistry Integrated dental floss and handles			
ISO 28319	2010-05	Ν	Dentistry Laser welding			
			Dentistry Products for external tooth			
ISO 28399	2011-01	N	bleaching			
			Medical devices Non-electrically driven			
ISO 28620	2010-02	N	portable infusion devices			
			Nanotechnologies Endotoxin test on			
			nanomaterial samples for in vitro			
			systems Limulus amebocyte lysate			
ISO 29701	2010-09	Ν	(LAL) test			
			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			
ISO 29781	2008-12	N	lower limb segment(s) present at birth			
			Dreath and anth and a factor is t			
			Prostheses and orthoses Factors to be			
180 20782	2008 12	N	considered when specifying a prosthesis for a person who has had a lower limb amputation			
ISO 29782	2008-12	N	person who has had a lower limb amputation			

			Prosthetics and orthotics Vocabulary		
ISO 29783-1	2008-12	Ν	Part 1: Normal gait		
130 29703-1	2000-12	IN	Fait_1. Normai gait		
			Condoms - Determination of nitrosamines		
ISO 29941	2010-12	Ν	migrating from natural rubber latex condoms		
100 200 11	2010 12		Prophylactic dams Requirements and test		
ISO 29942	2011-07	Ν	methods		
			Dentistry Zinc oxide/eugenol cements and		
ISO 3107	2011-03	Ν	zinc oxide/non-eugenol cements		
			Gas cylinders for medical use; Marking		
ISO 32	1977-05	N	for identification of content		
			Dentistry Root-canal instruments Part_1:		
ISO 3630-1	2008-02	N	General requirements and test methods		
			Dental root-canal instruments Part_2:		
ISO 3630-2	2000-12	N	Enlargers		 
100 0000 0	4004.00	N	Dental root-canal instruments; part_3:		
ISO 3630-3	1994-03	N	condensers, pluggers and spreaders Dentistry Root canal instruments Part_4:		 
ISO 3630-4	2009-07	Ν	Auxiliary instruments		
100 3030-4	2003-07	11	Dentistry Endodontic instruments Part_5:		
ISO 3630-5	2011-10	Ν	Shaping and cleaning instruments		
			Dental rotary instruments Burs Part_1:		
ISO 3823-1	1997-08	Ν	Steel and carbide burs		
			Dentistry Rotary bur instruments Part_2:		
ISO 3823-2	2003-05	Ν	Finishing burs		
			Dentistry Rotary bur instruments Part_2:		
ISO 3823-2 AMD 1	2008-07	N	Finishing burs; Amendment_1		
			Plastics collapsible containers for human		
			blood and blood components Part_1:		
ISO 3826-1	2003-11	N	Conventional containers		
			Plastics collapsible containers for human		
			blood and blood components Part_2:		
			Graphical symbols for use on labels and		
ISO 3826-2	2008-08	N	instruction leaflets		
			Plastics collapsible containers for human		
			blood and blood components Part_3:		
			Blood bag systems with integrated		
ISO 3826-3	2006-09	Ν	features		
			Acoustics Reference zero for the		
			calibration of audiometric equipment		
			Part_1: Reference equivalent threshold		
			sound pressure levels for pure tones and		
ISO 389-1	1998-11	Ν	supra-aural earphones		
			Acoustics Reference zero for the		
			calibration of audiometric equipment -		
			Part_2: Reference equivalent threshold		
			sound pressure levels for pure tones and		
ISO 389-2	1994-07	Ν	insert earphones		
00 000-2	1334-07	IN	insen earphones		 

			Acoustics - Reference zero for the				
			calibration of audiometric equipment				
			Part 3: Reference equivalent threshold				
			force levels for pure tones and bone				
ISO 389-3	1994-10	N	vibrators				
	100110		Acoustics - Reference zero for the				
			calibration of audiometric equipment				
			Part_3: Reference equivalent treshold				
			force levels for pure tones and bone				
ISO 389-3 Technical Corrige	1995-08	Ν	vibrators; Technical corrigendum_1				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_4: Reference levels for narrow-band				
ISO 389-4	1994-10	N	masking noise				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_6: Reference threshold of hearing				
ISO 389-6	2007-07	N	for test signals of short duration				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_7: Reference threshold of hearing				
			under free-field and diffuse-field listening				
ISO 389-7	2005-11	N	conditions				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_8: Reference equivalent threshold				
			sound pressure levels for pure tones and				
ISO 389-8	2004-05	N	circumaural earphones				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_9: Preferred test conditions for the				
			determination of reference hearing				
ISO 389-9	2009-05	N	threshold levels				
			Dentistry Designation system for teeth and				
ISO 3950	2009-05	N	areas of the oral cavity				
ISO 3964	1982-12	N	Dental handpieces; Coupling dimensions				
ISO 4049	2009-10	Ν	Dentistry Polymer-based restorative materials				
130 4049	2009-10	IN	Dentistry Information system on the location				
			of dental equipment in the working area of the				
ISO 4073	2009-07	Ν	oral health care provider				
			Natural latex rubber condoms Requirements				
ISO 4074	2002-02	N	and test methods				
			Network letter with an endowed D				
ISO 4074 Technical Corrigond	2002 10	Ν	Natural latex rubber condoms Requirements				
ISO 4074 Technical Corrigendu	2003-10	IN	and test methods; Technical Corrigendum_1	L	ļ	L	

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			Natural latex rubber condoms Requirements			
ISO 4074 Technical Corrigendu	2008-04	Ν	and test methods; Technical Corrigendum_2			
			Anaesthetic and respiratory equipment			
ISO 4135	2001-08	Ν	Vocabulary			
	2001.00					
ISO 4823	2000-12	Ν	Dentistry Elastomeric impression materials			
			Dentistry Elastomeric impression materials;			
ISO 4823 AMD 1	2007-07	N	Amendment_1			
			Dentistry Elastomeric impression materials;			
ISO 4823 Technical Corrigendu	2004-07	N	Technical Corrigendum_1			
			Anaesthetic and respiratory equipment			
			Conical connectors Part_1: Cones and			
ISO 5356-1	2004-05	N	sockets			
			Anaesthetic and respiratory equipment			
100 5050 0			Conical connectors Part_2: Screw-			
ISO 5356-2	2006-09	N	threaded weight-bearing connectors			
100 5359	1000.01	N	Anaesthetic machines for use with			
ISO 5358	1992-01	N	humans			
100 5050			Low-pressure hose assemblies for use			
ISO 5359	2008-06	N	with medical gases			
	0011 10	N	Low-pressure hose assemblies for use			
ISO 5359 AMD 1	2011-12	N	with medical gases; Amendment_1			
100 5000	0040.04	N	Anaesthetic vaporizers Agent-specific			
ISO 5360	2012-01	N	filling systems			
180 5201	1999-09	N	Anaesthetic and respiratory equipment			
ISO 5361 ISO 5361-4	1999-09	N N	Tracheal tubes and connectors Tracheal tubes; Part 4 : Cole type			
ISO 5362	2006-06	N	Anaesthetic reservoir bags			
ISO 5364	2000.07	N	Anaesthetic and respiratory equipment			
150 5364	2008-07	IN	Oropharyngeal airways			
			A possible tis and reprint on a subment			
			Anaesthetic and respiratory equipment			
ISO 5366-1	2000-12	N	Tracheostomy tubes Part_1: Tubes			
150 5366-1	2000-12	IN	and connectors for use in adults			
			Anaesthetic and respiratory equipment Tracheostomy tubes Part_3: Paediatric			
150 5266 2	2001-08	N	-			
ISO 5366-3	2001-08	N	tracheostomy tubes Anaesthetic and respiratory equipment		 	
			Tracheostomy tubes Part_3: Paediatric tracheostomy tubes; Technical			
ISO 5366-3 Technical Corrig	2002.01	N				
130 530-3 Technical Corrig	2003-01	N	Corrigendum_1 Breathing tubes intended for use with			
ISO 5367	2000-06	N	anaesthetic apparatus and ventilators			
190 3307	2000-00	IN	anaesmetic apparatus and ventilators			

			Implants for surgery Metallic materials		
ISO 5832-1	2007-06	Ν	Part_1: Wrought stainless steel		
100 3032-1	2007-00	IN	Implants for surgery Metallic materials		
			Part_1: Wrought stainless steel;		
ISO 5832-1 Technical	Corria 2008-04	Ν	Technical Corrigendum_1		
			Implants for surgery Metallic materials		
			Part_11: Wrought titanium 6-aluminium 7-		
ISO 5832-11	1994-09	Ν	niobium alloy		
			Implants for surgery Metallic materials		
			Part_12: Wrought cobalt-chromium-		
ISO 5832-12	2007-05	N	molybdenum alloy		
			Implants for surgery Metallic materials		
			Part_12: Wrought cobalt-chromium-		
			molybdenum alloy; Technical		
ISO 5832-12 Technica	al Corri 2008-09	N	Corrigendum_1		
			Implants for surgery Metallic materials		
			Part_14: Wrought titanium 15-		
			molybdenum 5-zirconium 3-aluminium		
ISO 5832-14	2007-10	N	alloy		 
100 5000 0	1000.07		Implants for surgery Metallic materials		
ISO 5832-2	1999-07	N	Part_2: Unalloyed titanium		
			Implants for surgery Metallic materials		
ISO 5832-3	1996-07	Ν	Part_3: Wrought titanium 6-aluminium 4- vanadium allov		
150 5652-5	1990-07	IN	Implants for surgery Metallic materials		
			Part_4: Cobalt-chromium-molybdenum		
ISO 5832-4	1996-07	Ν	casting alloy		
100 3032-4	1550-07	IN	Implants for surgery Metallic materials		
			Part_5: Wrought cobalt-chromium-		
ISO 5832-5	2005-10	Ν	tungsten-nickel alloy		
			Implants for surgery Metallic materials		
			Part_6: Wrought cobalt-nickel-chromium-		
ISO 5832-6	1997-07	Ν	molybdenum alloy		
			Implants for surgery; metallic materials;		
			part_7: forgeable and cold-formed cobalt-		
ISO 5832-7	1994-02	N	chromium-nickel-molybdenum-iron alloy		
			Implants for surgery Metallic materials		
			Part_8: Wrought cobalt-nickel-chromium-		
ISO 5832-8	1997-07	N	molybdenum-tungsten-iron alloy		 
			Implants for surgery Metallic materials		
	0007.00		Part_9: Wrought high nitrogen stainless		
ISO 5832-9	2007-06	N	steel		

			Implants for surgery Acrylic resin		
ISO 5833	2002-05	Ν	cements		
100 0000	2002.00		Implants for surgery Ultra-high-		
			molecular-weight polyethylene Part_1:		
ISO 5834-1	2005-06	Ν	Powder form		
			Implants for surgery Ultra-high-		
			molecular-weight polyethylene Part_1:		
ISO 5834-1 Technica	al Corrig 2007-05	Ν	Powder form; Technical Corrigendum_1		
			Implants for surgery Ultra-high-		
			molecular-weight polyethylene Part_2:		
ISO 5834-2	2011-08	N	Moulded forms		
			Implants for surgery Ultra-high-		
			molecular-weight polyethylene Part_3:		
ISO 5834-3	2005-07	N	Accelerated ageing methods		
			Implants for surgery Ultra-high-		
100 500 4 4			molecular-weight polyethylene Part_4:		
ISO 5834-4	2005-05	N	Oxidation index measurement method		
			Implants for surgery Ultra-high-		
	0005.00	N	molecular-weight polyethylene Part_5:		
ISO 5834-5	2005-06	N	Morphology assessment method		
			Implanta far aurganu matal hana aaraura		
			Implants for surgery; metal bone screws with hexagonal drive connection,		
			spherical under-surface of head,		
ISO 5835	1991-01	Ν	asymmetrical thread; dimensions		
130 3635	1991-01	IN	Implants for surgery; metal bone plates;		
			holes corresponding to screws with		
			asymmetrical thread and spherical under-		
ISO 5836	1988-12	Ν	surface		
100 0000	1000 12		Implants for surgery; Intramedullary		
			nailing systems; Part 1 : Intramedullary		
			nails with cloverleaf or V-shaped cross-		
ISO 5837-1	1985-06	Ν	section		
			Implants for surgery; Intramedullary		
ISO 5837-2	1980-11	Ν	nailing systems; Part 2 : Medullary pins		
			Implants for surgery Skeletal pins and		
			wires Part_1: Material and mechanical		
ISO 5838-1	1995-11	Ν	requirements		
			Implants for surgery; skeletal pins and		
			wires; part_2: Steinmann skeletal pins;		
ISO 5838-2	1991-01	N	dimensions	 	
			Implants for surgery; skeletal pins and		
ISO 5838-3	1993-09	N	wires; part_3: Kirschner skeletal wires		

			Cardiovascular implants Cardiac valve				
ISO 5840	2005-03	Ν	prostheses	Y	N		
			Implants for surgery Cardiac				
			pacemakers Part_2: Reporting of				
			clinical performance of populations of				
ISO 5841-2	2000-10	Ν	pulse generators or leads				
			Implants for surgery Cardiac				
			pacemakers Part_3: Low-profile				
			connectors [IS-1] for implantable				
ISO 5841-3	2000-10	Ν	pacemakers				
			Implants for surgery Cardiac				
			pacemakers Part_3: Low-profile				
			connectors (IS-1) for implantable				
ISO 5841-3 Technica	al Corrig 2003-11	Ν	pacemakers; Technical Corrigendum_1				
	<u> </u>		Conical fittings with a 6 % (Luer) taper for				
			syringes, needles and certain other				
			medical equipment; Part 1 : General				
ISO 594-1	1986-06	Ν	requirements				
			Conical fittings with 6%_(Luer) taper for				
			syringes, needles and certain other				
ISO 594-2	1998-09	Ν	medical equipment Part_2: Lock fittings				
			Reusable all-glass or metal-and-glass				
			syringes for medical use; Part 1 :				
ISO 595-1	1986-12	Ν	Dimensions				
			Reusable all-glass or metal-and-glass				
			syringes for medical use; Part 2 : Design,				
ISO 595-2	1987-12	Ν	performance requirements and tests				
			Hypodermic needles for single use;				
ISO 6009	1992-12	Ν	colour coding for identification				
			Hypodermic needles for single use				
			Colour coding for identification; Technical				
ISO 6009 Technical (	Corrigen 2008-03	Ν	Corrigendum_1				
			Dentistry Number coding system for rotary				
ISO 6360-1	2004-04	N	instruments Part_1: General characteristics			 	
			Dentistry Number coding system for rotary instruments Part_1: General characteristics;				
ISO 6360-1 Technical C	Corrigend 2007-09	Ν	Technical Corrigendum_1				
	50mgenu2007-03	IN	Dentistry Number coding system for rotary				
ISO 6360-2	2004-11	Ν	instruments Part_2: Shapes				
			Dentistry Number coding system for rotary				
ISO 6360-2 AMD 1	2011-12	N	instruments Part_2: Shapes; Amendment_1				

			Dentistry Number coding system for rotary			
			instruments - Part 3: Specific characteristics			
ISO 6360-3	2005-11	Ν	of burs and cutters			
130 0300-3	2003-11	IN	Dentistry Number coding system for rotary			
			instruments - Part 4: Specific characteristics			
ISO 6360-4	2004-06	Ν	of diamond instruments			
100 0000 4	2004 00	11	Dentistry Number coding system for rotary			
			instruments Part_5: Specific characteristics			
ISO 6360-5	2007-12	Ν	of root-canal instruments			
	2001 12		Dentistry Number coding system for rotary			
			instruments Part_6: Specific characteristics			
ISO 6360-6	2004-06	Ν	of abrasive instruments			
			Dentistry Number coding system for rotary			
			instruments Part_7: Specific characteristics			
ISO 6360-7	2006-02	Ν	of mandrels and special instruments			
			Implants for surgery Ceramic			
			materials Part_1: Ceramic materials			
ISO 6474-1	2010-02	Ν	based on high purity alumina			
			Implants for surgery; metal bone screws			
			with asymmetrical thread and spherical			
			under-surface; mechanical requirements			
ISO 6475	1989-11	Ν	and test methods			
130 0473	1909-11	IN	Single-use containers for venous blood		 -	
ISO 6710	1995-08	Ν	specimen collection			
ISO 6872	2008-09	N	Dentistry - Ceramic materials			
ISO 6873	1998-03	N	Dental gypsum products			
100 0075	1330-03	IN	Dentistry - Polymer-based pit and fissure			
ISO 6874	2005-08	Ν	sealants			
ISO 6875	2011-07	N	Dentistry Patient chair			
ISO 6876	2001-08	N	Dental root canal sealing materials			
ISO 6877	2001-08	N	Dentistry Root-canal obturating points			
130 0877	2000-04	IN	Surgical instruments; non-cutting,		 -	
	1000.10		articulated instruments; general			
ISO 7151	1988-12	N	requirements and test methods			
			Surgical instruments; metallic materials;			
ISO 7153-1	1991-04	N	part_1: stainless steel			
			Surgical instruments Metallic			
			materials Part_1: Stainless steel;			
ISO 7153-1 AMD 1	1999-03	Ν	Amendment_1			
			Wheelchairs Part_1: Determination of static			
ISO 7176-1	1999-10	N	stability			
			Wheelchairs Part_10: Determination of			
00			obstacle-climbing ability of electrically powered			
ISO 7176-10	2008-11	N	wheelchairs			
ISO 7176-11	1992-05	N	Wheelchairs; part_11: test dummies			
00 7/70 /0	1000		Wheelchairs; part_13: determination of			
ISO 7176-13	1989-08	N	coefficient of friction of test surfaces			

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			Wheelchairs Part_14: Power and control				
			systems for electrically powered wheelchairs				
100 7470 44	0000.00		and scooters Requirements and test				
ISO 7176-14	2008-02	N	methods				
			Wheelchairs Part_15: Requirements for				
100 7470 45	1000.11		information disclosure, documentation and				
ISO 7176-15	1996-11	N	labelling				
			Wheelchairs Part_16: Resistance to ignition				
100 7470 40	1007.05		of upholstered parts Requirements and test				
ISO 7176-16	1997-05	N	methods Wheelchairs Part_19: Wheeled mobility				
100 7476 40	2000.07	N	devices for use as seats in motor vehicles				
ISO 7176-19	2008-07	N	Wheelchairs Part_2: Determination of				
ISO 7176-2	2001-06	Ν	dynamic stability of electric wheelchairs				
150 / 176-2	2001-06	IN					
			Wheelchairs Part_21: Requirements and				
			test methods for electromagnetic compatibility				
			of electrically powered wheelchairs and				
ISO 7176-21	2009-04	Ν	scooters, and battery chargers				
ISO 7176-22	2000-05	N	Wheelchairs Part_22: Set-up procedures				
130 / 170-22	2000-03	IN	Wheelchairs - Part 23: Requirements and				
			test methods for attendant-operated stair-				
ISO 7176-23	2002-07	Ν	climbing devices				
130 / 170-23	2002-07	IN	Wheelchairs Part_24: Requirements and				
			test methods for user-operated stair-climbing				
ISO 7176-24	2004-10	Ν	devices				
ISO 7176-24	2007-04	N	Wheelchairs Part_26: Vocabulary				
130 / 176-20	2007-04	IN	Wheelchairs - Part 3: Determination of		-		
ISO 7176-3	2003-04	Ν	effectiveness of brakes				
1307170-3	2003-04	IN	Wheelchairs Part_4: Energy consumption of				
			electric wheelchairs and scooters for				
ISO 7176-4	2008-10	Ν	determination of theoretical distance range				
100 / 170 4	2000 10	11	Wheelchairs Part_5: Determination of				
ISO 7176-5	2008-06	Ν	dimensions, mass and manoeuvring space				
100 / 110 0	2000 00		Wheelchairs Part_6: Determination of				
			maximum speed, acceleration and				
ISO 7176-6	2001-10	Ν	deceleration of electric wheelchairs				
			Wheelchairs Part_7: Measurement of				
ISO 7176-7	1998-05	Ν	seating and wheel dimensions				
			Wheelchairs Part_8: Requirements and test				
			methods for static, impact and fatigue				
ISO 7176-8	1998-07	Ν	strengths				
			Wheelchairs Part_9: Climatic tests for				
ISO 7176-9	2009-11	Ν	electric wheelchairs				
ISO 7193	1985-12	Ν	Wheelchairs; Maximum overall dimensions				
			Neurosurgical implants Sterile, single-				
			use hydrocephalus shunts and				
ISO 7197	2006-06	N	components				
100 / 13/	2000-00	I N		Į	_ <u> </u>	1	<u> </u>

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			Neurosurgical implants Sterile, single-			
			use hydrocephalus shunts and			
ISO 7197 Technical Co	rrigen 2007-07	Ν	components; Technical Corrigendum_1			
			Cardiovascular implants - Tubular			
ISO 7198	1998-08	Ν	vascular prostheses			
100 / 130	1330-00		Cardiovascular implants and artificial			
			organs Blood-gas exchangers			
ISO 7199	2009-04	Ν	(oxygenators)			
100 / 100	2000 04		Cardiovascular implants and artificial			
			organs Blood-gas exchangers			
			(oxygenators) Amendment_1:			
			Clarifications for test methodologies,			
ISO 7199 AMD 1	2012-02	Ν	labelling, and sampling schedule			
100710071001	2012 02		Implants for surgery Partial and total			
			hip joint prostheses Part_1:			
			Classification and designation of			
ISO 7206-1	2008-04	Ν	dimensions			
	2000 01		Implants for surgery Partial and total			
			hip-joint prostheses Part_10:			
			Determination of resistance to static load			
ISO 7206-10	2003-12	Ν	of modular femoral heads			
100 1200 10	2000 12		Implants for surgery Partial and total			
			hip joint prostheses Part_2: Articulating			
			surfaces made of metallic, ceramic and			
ISO 7206-2	2011-04	Ν	plastics materials			
			Implants for surgery Partial and total			
			hip joint prostheses Part_4:			
			Determination of endurance properties			
			and performance of stemmed femoral			
ISO 7206-4	2010-06	Ν	components			
			Implants for surgery; partial and total hip			
			joint prostheses; part_6: determination of			
			endurance properties of head and neck			
ISO 7206-6	1992-03	Ν	region of stemmed femoral components			
			Implants for surgery Components for			
			partial and total knee joint prostheses			
			Part_1: Classification, definitions and			
ISO 7207-1	2007-02	Ν	designation of dimensions			
			Implants for surgery Components for			
			partial and total knee joint prostheses			
			Part_2: Articulating surfaces made of			
ISO 7207-2	2011-07	Ν	metal, ceramic and plastics materials			

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			Anaesthetic and respiratory equipment				
ISO 7376	2009-08	Ν	Laryngoscopes for tracheal intubation				
130 7370	2009-06	IN					
			Medical realized evotome				
			Medical gas pipeline systems Part_1:				
			Pipeline systems for compressed medical				
ISO 7396-1	2007-04	N	gases and vacuum				
			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				
ISO 7396-1 AMD 1	2010-01	Ν	hoses				
100 7 3 30 - 1 AMD 1	2010-01	IN	10363				
			Medical gas pipeline systems - Part 1:				
	0040.00		Pipeline systems for compressed medical				
ISO 7396-1 AMD 2	2010-02	N	gases and vacuum; Amendment_2				
			Medical gas pipeline systems Part_2:				
			Anaesthetic gas scavenging disposal				
ISO 7396-2	2007-04	N	systems				
			Dentistry Evaluation of biocompatibility of				
ISO 7405	2008-12	N	medical devices used in dentistry				
100 7400	0011.00	N	Copper-bearing contraceptive intrauterine				
ISO 7439	2011-06	N	devices Requirements and tests				
ISO 7488	1991-06	N	Dental amalgamators Dental materials - Determination of colour				
180 7404	2000.00	N	stability				
ISO 7491	2000-09	N					
ISO 7492	1997-02	N	Dental explorers				
ISO 7493	2006-05	N	Dentistry Operator's stool Dentistry Dental units Part_1: General				
ISO 7494-1	2011-08	Ν	requirements and test methods				
130 7494-1	2011-00	IN	Dentistry Dental units Part_2: Water and				
ISO 7494-2	2003-03	Ν	air supply				
ISO 7551	1996-12	N	Dental absorbent points				
100 1001	1000 12		Dental rotary instruments Diamond				
			instruments - Part 1: Dimensions,				
ISO 7711-1	1997-02	Ν	requirements, marking and packaging				
			Dental rotary instruments Diamond			1	
			instruments Part_1: Dimensions,			1	
			requirements, marking and packaging;			1	
ISO 7711-1 AMD 1	2009-05	Ν	Amendment_1			1	
			Dentistry Rotary diamond instruments				
ISO 7711-2	2011-07	N	Part_2: Discs				
			Dentistry Diamond rotary instruments				
			Part_3: Grit sizes, designation and colour				
ISO 7711-3	2004-11	N	code				

	1		Instruments for surgery; Scalpels with		T		
100 7740	1005 10	N					
ISO 7740	1985-12	N	detachable blades; Fitting dimensions				
			Instruments for surgery; Scissors and				
			shears; General requirements and test				
ISO 7741	1986-02	N	methods				
			Dental handpieces Part_1: High-speed air				
ISO 7785-1	1997-08	N	turbine handpieces				
100 7705 0	1005.00		Dental handpieces Part_2: Straight and				
ISO 7785-2	1995-08	N	geared angle handpieces Dental rotary instruments Laboratory				
ISO 7786	2001-04	N	abrasive instruments				
130 7780	2001-04	IN	Dental rotary instruments; Cutters; Part 1 :			+	
ISO 7787-1	1984-12	N	Steel laboratory cutters				
	1304-12	11	Dental rotary instruments Cutters Part_2:				
ISO 7787-2	2000-12	N	Carbide laboratory cutters				
	2000 12						
			Dental rotary instruments; cutters; part_3:				
ISO 7787-3	1991-12	N	carbide laboratory cutters for milling machines				
			Dental rotary instruments Cutters Part_4:				
ISO 7787-4	2002-03	N	Miniature carbide laboratory cutters				
ISO 7864	1993-05	N	Sterile hypodermic needles for single use				
			Dentistry Sterile injection needles for				
ISO 7885	2010-02	N	single use				
			Sterile hypodermic syringes for single				
ISO 7886-1	1993-10	N	use; part_1: syringes for manual use				
			Sterile hypodermic syringes for single				
			use Part_1: Syringes for manual use;				
ISO 7886-1 Technical Corri	1995-11	N	Technical Corrigendum 1				
	9 1000 11		Sterile hypodermic syringes for single				
			use Part_2: Syringes for use with				
ISO 7886-2	1996-05	Ν	power-driven syringe pumps				
100 1000-2	1330-03	IN IN	Sterile hypodermic syringes for single				
			use - Part 3: Auto-disable syringes for				
ISO 7886-3	2005-03	N	fixed-dose immunization				
130 7880-3	2005-05	IN				+	
			Sterile hypodermic syringes for single				
100 7000 4	0000 40		use Part_4: Syringes with re-use				
ISO 7886-4	2006-10	N	prevention feature		-		
	1000 55		Optics and optical instruments				
ISO 7944	1998-06	N	Reference wavelengths				
			Optics and optical instruments				
			Reference wavelengths; Technical				
ISO 7944 Technical Corrige	n 2009-07	N	Corrigendum_1				
			Ophthalmic optics Spectacle frames				
ISO 7998	2005-10	N	Lists of equivalent terms and vocabulary				

<b></b>			Mechanical contraceptives Reusable natural		1	
			and silicone rubber contraceptive			
ISO 8009	2004-10	Ν	diaphragms Requirements and tests			
	200110		Mechanical contraceptives Reusable natural			
			and silicone rubber contraceptive			
			diaphragms Requirements and tests;			
ISO 8009 AMD 1	2012-02	Ν	Amendment_1			
			Small-bore connectors for liquids and			
			gases in healthcare applications			
ISO 80369-1	2010-12	Ν	Part_1: General requirements			
			Medical electrical equipment Part_2-12:			
			Particular requirements for basic safety			
			and essential performance of critical care			
ISO 80601-2-12	2011-04	Ν	ventilators			
	2011 01					
			Medical electrical equipment Part_2-12:			
			Particular requirements for basic safety			
			and essential performance of critical care			
ISO 80601-2-12 Techn	ical C 2011-10	Ν	ventilators; Technical Corrigendum_1			
100 00001-2-12 160111			Medical electrical equipment Part_2-13:			
			Particular requirements for basic safety			
			and essential performance of an			
ISO 80601-2-13	2011-08	Ν	anaesthetic workstation			
130 80601-2-13	2011-00	IN				
			Medical electrical equipment Part_2-55:			
			Particular requirements for the basic			
ISO 80601-2-55	2011-12	N	safety and essential performance of			
150 80601-2-55	2011-12	N	respiratory gas monitors Medical electrical equipment Part_2-56:			
			Particular requirements for basic safety and			
			essential performance of clinical			
			thermometers for body temperature			
ISO 80601-2-56	2009-10	Ν	measurement			
			Medical electrical equipment Part_2-61:			
			Particular requirements for basic safety			
			and essential performance of pulse			
ISO 80601-2-61	2011-04	Ν	oximeter equipment			
			Non-invasive sphygmomanometers			
			Part_1: Requirements and test methods			
ISO 81060-1	2007-12	Ν	for non-automated measurement type			
			Non-invasive sphygmomanometers			
			Part_2: Clinical validation of automated			
ISO 81060-2	2009-05	Ν	measurement type			
			Non-invasive sphygmomanometers			
			Part_2: Clinical validation of automated			
			measurement type; Technical			
ISO 81060-2 Technical	Corri 2011-02	Ν	Corrigendum_1			
	001112011-02	IN	oongendum_r			

			Respiratory tract humidifiers for medical		Γ	[		Γ
			use Particular requirements for					
ISO 8185	2007-07	Ν	respiratory humidification systems					
150 0105	2007-07	IN	Radiation protection; Clothing for protection	· · · · · · · · · · · · · · · · · · ·				
			against radioactive contamination; Design,					
ISO 8194	1987-06	Ν	selection, testing and use					
			Acoustics Audiometric test methods					
			Part_1: Pure-tone air and bone					
ISO 8253-1	2010-11	N	conduction audiometry					
			Acoustics - Audiometric test methods -					
			Part 2: Sound field audiometry with pure-					
ISO 8253-2	2009-12	Ν	tone and narrow-band test signals					
100 0200 2	2000 12		Acoustics - Audiometric test methods -					
ISO 8253-3	2012-03	Ν	Part_3: Speech audiometry					
			Dental equipment Mercury and alloy mixers					
ISO 8282	1994-10	N	and dispensers					
			Orthopaedic instruments Drive					
			connections Part_1: Keys for use with					
ISO 8319-1	1996-05	Ν	screws with hexagon socket heads					
100 0010-1	1550-05		Orthopaedic instruments; Drive					
			connections; Part 2 : Screwdrivers for					
			single slot head screws, screws with					
			cruciate slot and cross-recessed head					
ISO 8319-2	1986-10	Ν	screws					
			Dentistry Test methods for rotary					
ISO 8325	2004-09	N	instruments					
10.0 00.50	1000.10		Oxygen concentrators for medical use					
ISO 8359	1996-12	N	Safety requirements					
			Injection containers and accessories					
ISO 8362-1	2009-12	Ν	Part_1: Injection vials made of glass tubing					
150 0502-1	2003-12	IN	Injection containers and accessories					
ISO 8362-2	2008-10	Ν	Part_2: Closures for injection vials					
100 0002 2	2000 10							
			Injection containers and accessories					
ISO 8362-3	2001-12	Ν	Part_3: Aluminium caps for injection vials					
			Injection containers and accessories					
			Part_4: Injection vials made of moulded					
ISO 8362-4	2011-09	N	glass					
			Injection containers and accessories					
100 0000 5			Part_5: Freeze drying closures for					
ISO 8362-5	2008-10	N	injection vials				-	-
			Injection containers and accessories					
	2010.00	NI	Part_6: Caps made of aluminium-plastics					
ISO 8362-6	2010-06	N	combinations for injection vials					

	<u> </u>		Injection containers and accessories	1	I	
			Part_7: Injection caps made of aluminium-			
ISO 8362-7	2000 04	N	plastics combinations without overlapping			
150 8362-7	2006-04	Ν	plastics part			
100 0 100	4000.00		Optics and optical instruments;			
ISO 8429	1986-09	N	Ophthalmology; Graduated dial scale			
100.0500.4			Infusion equipment for medical use			
ISO 8536-1	2011-09	N	Part_1: Infusion glass bottles			
			Infusion equipment for medical use			
			Part_10: Accessories for fluid lines for			
ISO 8536-10	2004-10	N	use with pressure infusion equipment			
			Infusion equipment for medical use			
			Part_11: Infusion filters for use with			
ISO 8536-11	2004-10	N	pressure infusion equipment			
			Infusion equipment for medical use			
ISO 8536-12	2007-04	N	Part_12: Check valves			
			Infusion equipment for medical use			
ISO 8536-2	2010-03	N	Part_2: Closures for infusion bottles			
			Infusion equipment for medical use			
			Part_3: Aluminium caps for infusion			
ISO 8536-3	2009-06	Ν	bottles			
			Infusion equipment for medical use			
			Part_4: Infusion sets for single use,			
ISO 8536-4	2010-10	Ν	gravity feed			
			Infusion equipment for medical use			
			Part_5: Burette infusion sets for single			
ISO 8536-5	2004-02	Ν	use, gravity feed			
			Infusion equipment for medical use			
			Part_6: Freeze drying closures for			
ISO 8536-6	2009-11	Ν	infusion bottles			
			Infusion equipment for medical use			
			Part_7: Caps made of aluminium-plastics			
ISO 8536-7	2009-01	Ν	combinations for infusion bottles			
			Infusion equipment for medical use			
			Part_8: Infusion equipment for use with			
ISO 8536-8	2004-08	Ν	pressure infusion apparatus			
	200+00		Infusion equipment for medical use			
			Part_9: Fluid lines for use with pressure			
ISO 8536-9	2004-10	Ν	infusion equipment			
8-0000	2004-10	IN	Sterile single-use syringes, with or			
ISO 8537	2007-10	Ν	without needle, for insulin			
130 0337	2007-10	IN				
			Prosthetics and orthotics: limb			
100 05 40 4	1000.00	NI	deficiencies; part_1: method of describing			
ISO 8548-1	1989-08	N	limb deficiencies present at birth			

	1	1				
			Prosthetics and orthotics; limb			
			deficiencies; part_2: method of describing			
ISO 8548-2	1993-07	N	lower limb amputation stumps			
			Prosthetics and orthotics; limb			
10.0 05 40 0			deficiencies; part_3: method of describing			
ISO 8548-3	1993-07	N	upper limb amputation stumps			
			Prosthetics and orthotics Limb			
			deficiencies Part_4: Description of			
ISO 8548-4	1998-07	N	causal conditions leading to amputation			
			Prosthetics and orthotics Limb			
1			deficiencies Part_5: Description of the			
			clinical condition of the person who has			
ISO 8548-5	2003-07	Ν	had an amputation			
			Prosthetics and orthotics; vocabulary; part_1:			
			general terms for external limb protheses and			
ISO 8549-1	1989-07	N	external orthoses			
			Prosthetics and orthotics; vocabulary; part 2:			
			terms relating to external limb prostheses and			
ISO 8549-2	1989-07	N	wearers of these prostheses			
			Prosthetics and orthotics; vocabulary; part_3:			
ISO 8549-3	1989-07	N	terms relating to external orthoses			
			Prosthetics and orthotics Functional			
			deficiencies Description of the person to be treated with an orthosis, clinical objectives of			
			treatment, and functional requirements of the			
ISO 8551	2003-08	Ν	orthosis			
			Ophthalmic optics Visual acuity			
			testing Standard optotype and its			
ISO 8596	2009-07	Ν	presentation			
			Optics and optical instruments			
ISO 8598	1996-08	N	Focimeters			
			Optics and optical instruments			
ISO 8598 Technical Corriger	1998-05	N	Focimeters; Technical corrigendum_1			
			Optics and photonics Medical			
			endoscopes and endotherapy devices			
ISO 8600-1	2005-05	Ν	Part_1: General requirements			
100 0000-1	2000-00	IN	Optics and optical instruments - Medical			
			endoscopes and endoscopic			
			accessories Part_2: Particular			
ISO 8600-2	2002-08	N	requirements for rigid bronchoscopes			
100 0000-2	2002-00	11	requirements for fight bronchoscopes	1	1	

			Optics and optical instruments Medical			
			endoscopes and endoscopic			
			accessories Part_3: Determination of			
100 0000 0	1007.07	N	field of view and direction of view of			
ISO 8600-3	1997-07	N	endoscopes with optics			
			Ontine and antine line to use ante. Madical			
			Optics and optical instruments Medical			
			endoscopes and endoscopic			
			accessories Part_3: Determination of			
	0000 40	N	field of view and direction of view of			
ISO 8600-3 AMD 1	2003-12	N	endoscopes with optics; Amendment_1			
l .			Optics and optical instruments Medical			
			endoscopes and certain accessories Part 4: Determination of maximum width			
ISO 8600-4	1997-07	Ν	of insertion portion			
150 8600-4	1997-07	IN				 
			Optics and photonics Medical			
			endoscopes and endotherapy devices			
			Part 5: Determination of optical			
ISO 8600-5	2005-03	N	resolution of rigid endoscopes with optics			
150 0000-5	2003-03	IN	Optics and photonics Medical			
			endoscopes and endotherapy devices			
ISO 8600-6	2005-03	Ν	Part_6: Vocabulary			
ISO 8612	2009-10	N	Ophthalmic instruments Tonometers			
100 0012	2000 10					
			Implants for surgery; fixation devices for			
ISO 8615	1991-11	Ν	use in the ends of the femur in adults			
			Ophthalmic optics Spectacle frames			
ISO 8624	2011-02	Ν	Measuring system and terminology			
			Cardiovascular implants and			
			extracorporeal systems			
			Haemodialysers, haemodiafilters,			
ISO 8637	2010-07	Ν	haemofilters and haemoconcentrators			
			Cardiovascular implants and			
			extracorporeal systems Extracorporeal			
			blood circuit for haemodialysers,			
ISO 8638	2010-07	Ν	haemodiafilters and haemofilters			
ISO 8669-1	1988-07	N	Urine collection bags; part_1: vocabulary			
			Urine collection bags Part_2: Requirements			
ISO 8669-2	1996-12	N	and test methods			
ISO 8670-1	1988-07	N	Ostomy collection bags; part_1: vocabulary			
100 0070 0	1006 10	N	Ostomy collection bags Part_2:			
ISO 8670-2	1996-12	N	Requirements and test methods		I	

[			Ostomy collection bags Part_3:	 	1	
			Determination of odour transmission of			
ISO 8670-3	2000-03	Ν	colostomy and ileostomy bags			
100 00/0 0	2000 00	N	Implants for surgery; staples with parallel			
			legs for orthopaedic use; general			
ISO 8827	1988-10	Ν	requirements			
100 0027	1900-10	IN	Implants for surgery; guidance on care			
ISO 8828	1988-10	N	and handling of orthopaedic implants			
150 0020	1900-10	IN	Inhalational anaesthesia systems			
			Part_7: Anaesthetic systems for use in			
			areas with limited logistical supplies of			
ISO 8835-7	2011-11	Ν	electricity and anaesthetic gases			
100 0000-1	2011-11	IN	Suction catheters for use in the			
ISO 8836	2007-09	Ν	respiratory tract			
150 0030	2007-03	IN				
			Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_1:			
ISO 8871-1	2003-10	Ν	Extractables in aqueous autoclavates			
100 007 1-1	2003-10					
			Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_2:			
ISO 8871-2	2003-10	Ν	Identification and characterization			
100 007 1 2	2000 10		Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_2:			
			Identification and characterization;			
ISO 8871-2 AMD 1	2005-07	Ν	Amendment_1			
	2000 01		, anonamona_			
			Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_3:			
ISO 8871-3	2003-08	Ν	Determination of released-particle count			
			Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_4:			
ISO 8871-4	2006-06	Ν	Biological requirements and test methods			
			Elastomeric parts for parenterals and for			
			devices for pharmaceutical use Part_5:			
ISO 8871-5	2005-08	Ν	Functional requirements and testing			
			Aluminium caps for transfusion, infusion			
			and injection bottles General			
ISO 8872	2003-03	Ν	requirements and test methods			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_1: Specifications			
ISO 8980-1	2004-02	Ν	for single-vision and multifocal lenses			

		1				
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_1: Specifications			
			for single-vision and multifocal lenses;			
ISO 8980-1 Technical Corric	2006-08	N	Technical Corrigendum_1			
	2000-00	IN IN				
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_2: Specifications			
ISO 8980-2	2004-02	N	for progressive power lenses			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_2: Specifications			
			for progressive power lenses; Technical			
ISO 8980-2 Technical Corrig	2006-08	N	Corrigendum_1			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_3: Transmittance			
ISO 8980-3	2003-10	N	specifications and test methods			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_4: Specifications			
			and test methods for anti-reflective			
ISO 8980-4	2006-08	N	coatings			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_5: Minimum			
			requirements for spectacle lens surfaces			
ISO 8980-5	2005-08	N	claimed to be abrasion-resistant			
			Dentistry Hose connectors for air driven			
ISO 9168	2009-07	N	dental handpieces			
			Terminal units for medical gas pipeline			
			systems Part_1: Terminal units for use			
100 0170 1	0000.07	N	with compressed medical gases and			
ISO 9170-1	2008-07	N	vacuum Terminal units for medical gas pipeline			
			systems - Part 2: Terminal units for			
ISO 9170-2	2008-07	N	anaesthetic gas scavenging systems			
130 9170-2	2006-07	IN	Dentistry - Extraction forceps - Part 1:			
ISO 9173-1	2006-06	N	General requirements and test methods			
			Dentistry Extraction forceps Part_2:			
ISO 9173-2	2010-05	N	Designation			
			Injection equipment for medical use			
ISO 9187-1	2010-10	N	Part_1: Ampoules for injectables			
			Injection equipment for medical use			
ISO 9187-2	2010-10	N	Part_2: One-point-cut (OPC) ampoules			
			Implants for surgery; metal bone screws			
			with conical under-surface of head;			
ISO 9268	1988-12	N	dimensions			

			Implants for surgery; metal bone plates;		
			holes and slots corresponding to screws		
ISO 9269	1988-12	Ν	with conical under-surface		
ISO 9333	2006-07	N	Dentistry Brazing materials		
			Optics and optical instruments Test		
			lenses for calibration of focimeters		
			Part_1: Test lenses for focimeters used		
ISO 9342-1	2005-05	Ν	for measuring spectacle lenses		
			Optics and optical instruments Test		
			lenses for calibration of focimeters		
			Part_2: Test lenses for focimeters used		
ISO 9342-2	2005-11	N	for measuring contact lenses		
			Anaesthetic and respiratory equipment		
			Heat and moisture exchangers (HMEs)		
			for humidifying respired gases in humans - Part 1: HMEs for use with		
ISO 9360-1	2000-03	Ν	minimum tidal volumes of 250 ml		
100 3300-1	2000-03	11			
			Anaesthetic and respiratory equipment		
			Heat and moisture exchangers (HMEs)		
			for humidifying respired gases in		
			humans Part_2: HMEs for use with		
			tracheostomized patients having		
ISO 9360-2	2001-04	N	minimum tidal volumes of 250_ml		
			Power-operated lifting platforms for persons		
			with impaired mobility Rules for safety,		
ISO 9386-1	2000-11	Ν	dimensions and functional operation Part_1: Vertical lifting platforms		
100 3300 1	2000 11	N			
			Power-operated lifting platforms for persons		
			with impaired mobility Rules for safety,		
			dimensions and functional operation Part_2:		
ISO 9386-2	2000-11	N	Powered stairlifts for seated, standing and wheelchair users moving in an inclined plane		
100 3300-2	2000-11	IN	Ophthalmic optics Contact lenses and		
			contact lens care products -		
			Determination of biocompatibility by		
ISO 9394	1998-08	Ν	ocular study with rabbit eyes		
			Implants for surgery; non-destructive		
			testing; liquid penetrant inspection of		
ISO 9583	1993-10	Ν	metallic surgical implants		
			Implants for surgery; non-destructive		
			testing; radiographic examination of cast		
ISO 9584	1993-10	Ν	metallic surgical implants		

			Implants for surgery; determination of			
100 0505	1000 10		bending strength and stiffness of bone			
ISO 9585	1990-12	N	plates			
			Stainless steel needle tubing for			
ISO 9626	1991-09	N	manufacture of medical devices			
			Stainless steel needle tubing for the			
			manufacture of medical devices;			
ISO 9626 AMD 1	2001-06	N	Amendment_1			
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			
	1000 12		Metal-ceramic dental restorative systems;			
ISO 9693 AMD 1	2005-10	N	Amendment 1			
	2000 10		Dentistry Compatibility testing Part_1:			
ISO 9693-1	2012-02	N	Metal-ceramic systems			
			Neurosurgical implants Self-closing			
ISO 9713	2002-09	N	intracranial aneurysm clips			
100 01 10	2002 00					
			Orthopaedic drilling instruments; part_1:			
ISO 9714-1	1991-03	N				
150 97 14-1	1991-03	N	drill bits, taps and countersink cutters			
			Ophthalmic instruments Trial case			
ISO 9801	2009-12	N	lenses			
10.0 0070			Dental hand instruments Reusable mirrors			
ISO 9873	1998-11	N	and handles			
			Dental hand instruments Reusable mirrors			
ISO 9873 Technical Corrigendu		Ν	and handles; Technical Corrigendum_1			
150 9873 Technical Comgendu	1 2000-06	IN	Dentistry Water-based cements Part_1:			
ISO 9917-1	2007-10	N	Powder/liquid acid-base cements			
130 9917-1	2007-10	IN	Dentistry Water-based cements Part_2:			
ISO 9917-2	2010-04	N	Resin-modified cements			
100 3311 2	2010 04		Urine absorbing aids; vocabulary; part_1:			
ISO 9949-1	1993-07	N	conditions of urinary incontinence			
			Urine absorbing aids; vocabulary; part 2:			
ISO 9949-2	1993-07	N	products			
			Urine absorbing aids; vocabulary; part_3:			
ISO 9949-3	1993-07	N	identification of product types			
ISO 9997	1999-12	N	Dental cartridge syringes			
			Assistive products for persons with disability			
ISO 9999	2011-07	N	Classification and terminology			
			Electronic Health Record-System Functional			
ISO/HL7 10781	2009-11	N	Model, Release_1.1			
			Health informatics HL_7 version_3			
ISO/HL7 21731	2006-08	N	Reference information model Release_1			
			Data Exchange Standards Health Level			
			Seven Version_2.5 An application protocol			
			for electronic data exchange in healthcare			
ISO/HL7 27931	2009-07	N	environments			
			Data Exchange Standards HL7 Clinical			
ISO/HL7 27932	2009-12	N	Document Architecture, Release_2			

			Health informatics Common terminology				
ISO/HL7 27951	2009-11	Ν	services, release_1	l l			
	2000		Health informatics Individual case safety				
			reports (ICSRs) in pharmacovigilance -	1			
			Part 1: Framework for adverse event	1			
ISO/HL7 27953-1	2011-12	N	reporting	1			
			Health informatics Individual case safety				
			reports (ICSRs) in pharmacovigilance	1			
			Part_2: Human pharmaceutical reporting	1			
ISO/HL7 27953-2	2011-12	N	requirements for ICSR	1			
			Information technology Office equipment				
			accessibility guidelines for elderly persons and	1			
ISO/IEC 10779	2008-06	N	persons with disabilities	1			
			Information technology Interoperability with				
			assistive technology (AT) Part_1:	1			
			Requirements and recommendations for	1			
ISO/IEC 13066-1	2011-05	Ν	interoperability	1			
			Information technology User interfaces	1			
ISO/IEC 29136	2012-05	Ν	Accessibility of personal computer hardware	1			
			Information technology Survey of icons and				
			symbols that provide access to functions and	1			
			facilities to improve the use of information	1			
			technology products by the elderly and	1			
ISO/IEC TR 19765	2007-07	N	persons with disabilities	1			
			Information technology Guidelines for the				
			design of icons and symbols accessible to all	1			
			users, including the elderly and persons with	1			
ISO/IEC TR 19766	2007-06	N	disabilities	1			
			Information technology Accessibility				
			considerations for people with disabilities	1			
ISO/IEC TR 29138-1	2009-06	N	Part_1: User needs summary	1			
			Information technology Accessibility				
			considerations for people with disabilities	1			
ISO/IEC TR 29138-2	2009-06	N	Part_2: Standards inventory	1			
			Information technology Accessibility				
			considerations for people with disabilities	1			
ISO/IEC TR 29138-3	2009-06	N	Part_3: Guidance on user needs mapping	1			
			Health informatics Point-of-care				
			medical device communication	1			
ISO/IEEE 11073-10101	2004-12	N	Part 10101: Nomenclature	1			
						1	
	1		Health informatics Point-of-care	1		1	
				1			
	0004.40	N	medical device communication	1			
ISO/IEEE 11073-10201	2004-12	N	Part_10201: Domain information model	·	+		
			Health informatics Personal health device	1			
	0040.05		communication Part_10404: Device	1			
ISO/IEEE 11073-10404	2010-05	N	specialization Pulse oximeter	·	+		
	1		Health informatics Personal health device	1		1	
	0040.05		communication Part_10407: Device	1			
ISO/IEEE 11073-10407	2010-05	N	specialization Blood pressure monitor	I			

			Health informatics Point-of-care				
			medical device communication -				
			Part_10408: Device specialization				
ISO/IEEE 11073-10408	2010-05	Ν	Thermometer				
130/IEEE 110/3-10400	2010-03	IN	Health informatics Point-of-care				
			medical device communication -				
			—				
ISO/IEEE 11073-10415	2010-05	Ν	Part_10415: Device specialization Weighing scale				
150/IEEE 11073-10415	2010-05	IN	Health informatics Personal health device				
			communication Part_10417: Device				
ISO/IEEE 11073-10417	2010-05	Ν	specialization Glucose meter				
	2010 00		Health informatics - Point-of-care				
			medical device communication -				
			Part_10471: Device specialization				
ISO/IEEE 11073-10471	2010-05	Ν	Independant living activity hub				
	2010-03	IN	Health informatics - Point-of care				
			medical device communications -				
			Part_20101: Application profiles; Base				
ISO/IEEE 11073-20101	2004-12	N	standard				
130/IEEE 110/3-20101	2004-12	IN	Health informatics Point-of-care				
			medical device communication				
			Part_20601: Application profile				
ISO/IEEE 11073-20601	2010-05	N					
150/IEEE 11073-20601	2010-05	N	Optimized exchange protocol				
			Health informatics Point-of-care				
			medical device communications				
	0004.40	NI	Part_30200: Transport profile; Cable				
ISO/IEEE 11073-30200	2004-12	N	connected				
			Health informatics Point-of-care				
			medical device communications				
	000440		Part_30300: Transport profile; Infrared				
ISO/IEEE 11073-30300	2004-12	N	wireless Dental implants; guidelines for developing				
ISO/TR 11175	1993-08	Ν	dental implants				
130/11/11/13	1993-00	IN	Health informatics Clinical stakeholder				
ISO/TR 11487	2008-12	Ν	participation in the work of ISO_TC 215				
			Communication aids for blind persons				
			Identifiers, names and assignation to coded				
			character sets for 8-dot Braille characters				
			Part_1: General guidelines for Braille				
ISO/TR 11548-1	2001-12	N	identifiers and shift marks				
			Communication aids for blind persons				
			Identifiers, names and assignation to coded character sets for 8-dot Braille characters -				
ISO/TR 11548-2	2001-12	N	Part_2: Latin alphabet based character sets				
100/11/11040-2	2001-12	IN	n an_2. Latin alphabet based chaldbler sets	1	1	1	

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			Health informatics Interoperability of			
			telehealth systems and networks - Part 1:			
ISO/TR 16056-1	2004-07	Ν	Introduction and definitions			
130/TR 16036-1	2004-07	IN	Health informatics - Interoperability of		-	
			telehealth systems and networks Part_2:			
ISO/TR 16056-2	2004-07	Ν	Real-time systems			
150/TR 10050-2	2004-07	IN	Real-lime systems			
			Medical devices Guidance on the			
			selection of standards in support of			
			recognized essential principles of safety			
ISO/TR 16142	2006-01	N	and performance of medical devices			
			Health informatics Health informatics			
ISO/TR 17119	2005-01	N	profiling framework			
			Clinical laboratory testing and in vitro			
			diagnostic test systems In vitro			
			diagnostic medical devices for			
			professional use Summary of			
			regulatory requirements for information			
ISO/TR 18112	2006-01	N	supplied by the manufacturer			
130/11/10112	2000-01	IN				
			Health informatics Interoperability and			
			compatibility in messaging and communication			
ISO/TR 18307	2001-12	Ν	standards Key characteristics			
	2001.12		Health informatics Electronic health record			
ISO/TR 20514	2005-10	Ν	Definition, scope and context			
			Ophthalmic instruments Background			
			for light hazard specification in			
ISO/TR 20824	2007-07	Ν	ophthalmic instrument standards			
100/11(20021	2001 01		Health informatics - Trusted end-to-end			
ISO/TR 21089	2004-06	Ν	information flows			
			Health informatics Security requirements for			
			archiving of electronic health records			
ISO/TR 21548	2010-02	Ν	Guidelines			
			Health informatics Use of mobile			
			wireless communication and computing			
			technology in healthcare facilities			
			Recommendations for electromagnetic			
			compatibility (management of			
	0007.00		unintentional electromagnetic			
ISO/TR 21730	2007-02	N	interference) with medical devices			
	0000.44	N	Health informatics Good principles and			
ISO/TR 22221	2006-11	N	practices for a clinical data warehouse			
			Ergonomics data and guidelines for the			
			Ergonomics data and guidelines for the application of ISO/IEC_Guide 71 to products			
			and services to address the needs of older			
ISO/TR 22411	2008-09	Ν	persons and persons with disabilities			
100/ 1 K 22411	2000-09	IN	persons and persons with disabilities	1	1	1

ISO/TR 22442-4	2010-12	Ν	Medical devices utilizing animal tissues and their derivatives Part_4: Principles for elimination and/or inactivation of transmissible spongiform encephalopathy (TSE) agents and validation assays for those processes			
ISO/TR 22676	2006-10	N	Prosthetics Testing of ankle-foot devices and foot units Guidance on the application of the test loading conditions of ISO_22675 and on the design of appropriate test equipment			
			Health informatics Functional characteristics			
ISO/TR 22790	2007-12	Ν	of prescriber support systems			
			Ophthalmic implants Intraocular lenses Guidance on assessment of the need for clinical investigation of			
ISO/TR 22979	2006-02	N	intraocular lens design modifications			
	0040.00		Cosmetics Good Manufacturing Practices			
ISO/TR 24475	2010-03	N	General training document Health informatics Business requirements			
ISO/TR 25257	2009-09	N	for an international coding system for medicinal products			
ISO/TR 27809	2007-07	N	Health informatics Measures for ensuring patient safety of health software			
ISO/TR 28642	2011-07	N	Dentistry Guidance on colour measurement			
ISO/TR 28980	2007-01	N	Ophthalmic optics Spectacle lenses Parameters affecting lens power measurement			
ISO/TR 9586	1988-12	N	Implants for surgery; usage of the terms "valgus" and "varus" in orthopaedic surgery			
		Standards total	1102			
			#DE71101			
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