Status: 2014-S	September				-	Brasil	_	
				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/REGIONAL REFERENCE NO.	PLEASE FILL IN:	PLEASE FILL IN:
Document Reference	Publication	Status N- Standard, N-E -Draft, VN-E predraft,	English Title	Recognised ? Y- fully, P-partial,N- NO		National Reference	Publication date of the national standard	Recognition Number, if available
IEC 60118-0	1983	N	Measurement of electroacoustical characteristics	Y	Y	There is no national reference		IN nº 09/13
IEC 60118-0 AM		N	Hearing aids; part_0: measurement of electroacoustical characteristics; amendment_1	Y	Y	There is no national reference		IN nº 09/13
IEC 60118-1	1995-04	N	Hearing aids Part_1: Hearing aids with induction pick-up coil input					
IEC 60118-1 AM	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 Edit		N	induction pick-up coil input Hearing aids Part_12: Dimensions of					
IEC 60118-12 IEC 60118-13	1996-09 2011-04	N N	electrical connector systems Electroacoustics Hearing aids Part_13: Electromagnetic compatibility (EMC)	Y	Y	There is no national reference		IN nº 09/13
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					
IEC 60118-2	1983	Ν	Hearing aids. Part 2 : Hearing aids with automatic gain control circuits					
IEC 60118-2 AM	1993-02	N	Hearing aids; part_2: hearing aids with automatic gain control circuits; amendment_1 Hearing aids Part_2: Hearing aids with					
IEC 60118-2 AM	1997-05	Ν	automatic gain control circuits; Amendment_2					

			Electropopulation I legitar side Dart 4.		L			
			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	N	earphones					
			Hearing aids Part_6: Characteristics of					
IEC 60118-6	1999-06	N	electrical input circuits for hearing aids					
			Electroacoustics Hearing aids Part_7:					
			Measurement of performance					
			characteristics of hearing aids for	Y	Y			
			production, supply and delivery quality			There is no national		
IEC 60118-7	2005-10	N	assurance purposes			reference		IN nº 09/13
			Electroacoustics Hearing aids Part_8:					
			Methods of measurement of performance					
			characteristics of hearing aids under					
IEC 60118-8	2005-10	N	simulated in situ working conditions					
			Hearing aids. Part 9 : Methods of					
			measurement of characteristics of hearing					
IEC 60118-9	1985	Ν	aids with bone vibrator output					
			Electroacoustics Simulators of human					
			head and ear Part_4: Occluded-ear					
			simulator for the measurement of earphones					
IEC 60318-4	2010-01	N	coupled to the ear by means of ear inserts					
			Household and similar electrical					
			appliances Safety Part_2-52: Particular					
IEC 60335-2-52	2005 10	N	requirements for oral hygiene appliances					
IEC 00335-2-52	2003-10	IN	Household and similar electrical					
			appliances Safety Part_2-52: Particular					
			requirements for oral hygiene appliances;					
IEC 60335-2-52	2008 04	N	Amendment 1					
IEC 00335-2-52	12000-04	IN	Amendment_1					
			Household and similar electrical					
			appliances Safety Part_2-52: Particular					
IEC 60335-2-52	2008-07	Ν	requirements for oral hygiene appliances					
120 00000 2 02	2000 01							
			Medical electrical equipment X-ray					
	0005.04		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 Co	ri 2006-05	N	Corrigendum_1					

			Determination of the permanent filtration					
IEC 60522	2003-12	N	of X-ray tube assemblies					
			High-voltage cable plug and socket					
	10-0		connections for medical X-ray					
IEC 60526	1978	N	equipment					
			High-voltage cable plug and socket					
			connections for medical X-ray					
IEC 60526 Co	ri 2010-04	N	equipment					
			Medical electrical equipment Dose					
IEC 60580	2003-09	N	area product meters					
			Madical electrical equipment Dart 1:					
			Medical electrical equipment Part_1: General requirements for basic safety	Y	Y	ABNT NBR IEC 60601-1	2010	IN nº 09/13
IEC 60601-1	2005-12	N	and essential performance					
IEC 00001-1	2005-12	IN	Medical electrical equipment Part_1:					
			General requirements for basic safety					
			and essential performance;	Y	Y	ABNT NBR IEC 60601-1	2010	IN nº 09/13
IEC 60601-1 C	2006-12	Ν	Corrigendum_1					
	2000-12		Medical electrical equipment Part_1:					
			General requirements for basic safety					
			and essential performance;	Y	Y	ABNT NBR IEC 60601-1	2010	IN nº 09/13
IEC 60601-1 C	2007-12	N	Corrigendum_2					
			Medical electrical equipment Part_1:				0010	
			General requirements for basic safety	Y	Y	ABNT NBR IEC 60601-1	2010	IN nº 09/13
IEC 60601-1 li	n 2008-04	Ν	and essential performance					
			Medical electrical equipment Part_1:					
			General requirements for basic safety	Y	Y	ABNT NBR IEC 60601-1	2010	IN nº 09/13
			and essential performance	I	T	ABINT INDR IEC 00001-1	2010	IN 11º 09/13
IEC 60601-1 li	n 2009-01	N	Interpretation sheet_2					
			Medical electrical equipment Part_1-					
			1: General requirements for safety;					
			Collateral standard: Safety	Y	Y	ABNT NBR IEC 60601-1-1	2004	IN nº 09/13
			requirements for medical electrical					
IEC 60601-1-1	2000-12	N	systems					
			Medical electrical equipment Part_1-					
			10: General requirements for basic					
			safety and essential performance	Y	Y	ABNT NBR IEC 60601-1-10	2010	IN nº 09/13
			Collateral Standard: Requirements for	•			2010	
			the development of physiologic closed-					
IEC 60601-1-1	(2007-11	N	loop controllers					

IEC 60601-1-1	12010-04	N	Medical electrical equipment Part_1- 11: General requirements for basic safety and essential performance Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	Y	Y	ABNT NBR IEC 60601-1-11	2012	IN nº 09/13
IEC 60601-1-1	2011-04	N	Medical electrical equipment Part_1- 11: General requirements for basic safety and essential performance Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment	Y	Y	ABNT NBR IEC 60601-1-11	2012	IN nº 09/13
IEC 60601-1-1		N	Medical electrical equipment Part_1- 11: General requirements for basic safety and essential performance Collateral standard: Requirements for medical electrical equipment and medical electrical systems used in the home healthcare environment; Technical Corrigendum_1	Y	Y	ABNT NBR IEC 60601-1-11	2012	IN nº 09/13
IEC 60601-1-2	2007-03	N	Medical electrical equipment Part_1- 2: General requirements for basic safety and essential performance Collateral standard: Electromagnetic compatibility Requirements and tests	Y	Y	ABNT NBR IEC 60601-1-2	2010	IN nº 09/13
IEC 60601-1-2	2010-03	N	Medical electrical equipment Part_1- 2: General requirements for basic safety and essential performance Collateral standard: Electromagnetic compatibility Requirements and tests	Y	Y	ABNT NBR IEC 60601-1-2	2010	IN nº 09/13
IEC 60601-1-3	2008-01	N	Medical electrical equipment Part_1-3: General requirements for basic safety and essential performance Collateral standard: Radiation protection in diagnostic X-ray equipment	Y	Y	ABNT NBR IEC 60601-1-3	2011	IN nº 09/13

				r		1		
IEC 60601-1-4	1996-05	N	Medical electrical equipment Part_1: General requirements for safety 4Collateral standard: Programmable electrical medical systems	Y	Y	ABNT NBR IEC 60601-1-4	2004	IN nº 09/13
IEC 60601-1-4 A	1999-10	Ν	Medical electrical equipment Part_1- 4: General requirements for safety Collateral standard: Programmable electrical medical systems; Amendment_1	Y	Y	ABNT NBR IEC 60601-1-4	2004	IN nº 09/13
IEC 60601-1-4 E	2000-04	N	Medical electrical equipment Part_1- 4: General requirements for safety Collateral standard: Programmable electrical medical systems	Y	Y	ABNT NBR IEC 60601-1-4	2004	IN nº 09/13
IEC 60601-1-6	2010-01	Ν	Medical electrical equipment General requirements for basic safety and essential performance Collateral Standard: Usability	Y	Y	ABNT NBR IEC 60601-1-6	2011	IN nº 09/13
IEC 60601-1-8	2006-10	Ν	Medical electrical equipment Part_1- 8: General requirements for basic safety and essential performance Collateral Standard: General requirements, tests and guidance for alarm systems in medical electrical equipment and medical electrical systems	Y	Y	ABNT NBR IEC 60601-1-8	2010	IN nº 09/13
IEC 60601-1-9	2007-07	N	Medical electrical equipment Part_1- 9: General requirements for basic safety and essential performance Collateral Standard: Requirements for environmentally conscious design	Y	Y	ABNT NBR IEC 60601-1-9	2010	IN nº 09/13
IEC 60601-2-1		N	Medical electrical equipment Part_2- 1: Particular requirements for the basic safety and essential performance of electron accelerators in the range 1_MeV to 50_MeV	Y	Y	ABNT NBR IEC 60601-2-1	2011	IN nº 09/13
IEC 60601-2-10		N	Medical electrical equipment; part_2: particular requirements for the safety of nerve and muscle stimulators	Ν	Ν			
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			Madical clastrical any instant Dart O		1	1	
			Medical electrical equipment Part_2-				
			10: Particular requirements for the	Ν	N		
			safety of nerve and muscle stimulators;				
IEC 60601-2-10	2001-09	N	Amendment_1				
			Medical electrical equipment Part_2-				
			10: Particular requirements for the	Ν	Ν		
			safety of nerve and muscle stimulators;				
IEC 60601-2-10	2002-02	N	Amendment_1				
			Medical electrical equipment Part_2:				
			Particular requirements for the safety of				
IEC 60601-2-11	1007.09	Ν	gamma beam therapy equipment				
IEC 00001-2-1	1997-00		Amendment_1 Medical electrical				
			equipment Part_2-11: Particular				
IEC 60601-2-11	2004 07	N	requirements for the safety of gamma beam therapy equipment				
IEC 00001-2-1	2004-07	N	Medical electrical equipment Part_2-				
			13: Particular requirements for the				
			safety and essential performance of				
IEC 60601-2-13	2002.05	Ν	anaesthetic systems				
IEC 00001-2-13	2003-05	IN	Medical electrical equipment Part_2-				
			13: Particular requirements for the				
			safety and essential performance of				
IEC 60601-2-13	2006 05	Ν	anaesthetic systems; Amendment_1				
	2000-03		Medical electrical equipment Part_2-				
			13: Particular requirements for the				
IEC 60601-2-13	2000-08	N	safety of anaesthetic systems				
	2009-00		Medical electrical equipment Part_2-				
			16: Particular requirements for basic				
			safety and essential performance of	Ν	Ν		
			haemodialysis, haemodiafiltration and		IN IN		
IEC 60601-2-16	2008-04	N	haemofiltration equipment				
	2000-04		Medical electrical equipment Part_2-				
			16: Particular requirements for basic				
			safety and essential performance of	Ν	Ν		
			haemodialysis, haemodiafiltration and	IN	IN		
IEC 60601-2-16	2008-10	N	haemofiltration equipment				
	2000-10	IN	Medical electrical equipment Part_2-				
			17: Particular requirements for the				
			safety of automatically-controlled				
IEC 60601-2-17	2005 00	N	brachytherapy afterloading equipment				
1EC 00001-2-17	2000-09	IN	practivitierapy attenoading equipment				

IEC 60601-2-1{2009-08	N	Medical electrical equipment Part_2- 18: Particular requirements for basic safety and essential performance of endoscopic equipment	Υ	Y	There is no national reference		IN nº 09/13
IEC 60601-2-1\$2009-02	N	Medical electrical equipment Part_2- 19: Particular requirements for the basic safety and essential performance of infant incubators	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-1\$2012-02	N	Medical electrical equipment Part_2- 19: Particular requirements for the basic safety and essential performance of infant incubators; Corrigendum_1					
IEC 60601-2-2 2009-02	N	Medical electrical equipment Part_2- 2: Particular requirements for the basic safety and essential performance of high frequency surgical equipment and high frequency surgical accessories	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-2(2009-02	N	Medical electrical equipment Part_2- 20: Particular requirements for the basic safety and essential performance of infant transport incubators	Y	Y	ABNT NBR IEC 60601-2-20	2012	IN nº 09/13
JEC 60601-2-2(2012-02	N	Medical electrical equipment Part_2- 20: Particular requirements for the basic safety and essential performance of infant transport incubators; Corrigendum_1	Y	Y	ABNT NBR IEC 60601-2-20	2012	IN nº 09/13
IEC 60601-2-212009-02	N	Medical electrical equipment Part_2- 21: Particular requirements for the basic safety and essential performance of infant radiant warmers	Y	Y	ABNT NBR IEC 60601-2-21	2013	IN nº 09/13
IEC 60601-2-242007-05	N	Medical electrical equipment Part_2- 22: Particular requirements for basic safety and essential performance of surgical, cosmetic, therapeutic and diagnostic laser equipment	Y	Y	ABNT NBR IEC 60601-2-22	2012	IN nº 09/13

IEC 60601-2-232011-02	N	Medical electrical equipment Part_2- 23: Particular requirements for the basic safety and essential performance of transcutaneous partial pressure monitoring equipment	Y	Y	ABNT NBR IEC 60601-2-23	2012	IN nº 09/13
IEC 60601-2-241998-02	N	Medical electrical equipment Part_2- 24: Particular requirements for the safety of infusion pumps and controllers	Ν	Ν			
IEC 60601-2-252011-10	N	Medical electrical equipment Part_2- 25: Particular requirements for basic safety and essential performance of electrocardiographs	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-262003-12	N	Medical electrical equipment Part_2- 26: Particular requirements for the safety of electroencephalographs	Ν	Ν			
IEC 60601-2-272011-03	N	Medical electrical equipment Part_2- 27: Particular requirements for the basic safety and essential performance of electrocardiographic monitoring equipment	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-2{2010-03	N	Medical electrical equipment Part_2- 28: Particular requirements for basic safety and essential performance of X- ray tube assemblies for medical diagnosis	Y	Y	ABNT NBR IEC 60601-2-28	2012	IN nº 09/13
IEC 60601-2-2§2008-06	N	Medical electrical equipment Part_2- 29: Particular requirements for the basic safety and essential performance of radiotherapy simulators					
IEC 60601-2-3 1991-06	N	Medical electrical equipment; part_2: particular requirements for the safety of short-wave therapy equipment	Ν	Ν			
IEC 60601-2-3 1998-09	N	Medical electrical equipment Part_2: Particular requirements for the safety of short-wave therapy equipment; Amendment_1	Υ	Y	ABNT NBR IEC 60601-2-3		IN nº 09/13

IN nº 09/13 IN nº 09/13					Medical electrical equipment Part_2-		
					31: Particular requirements for basic		
IN nº 09/13	2013	ABNT NBR IEC 60601-2-31	Y	Y	safety and essential performance of		
IN nº 09/13					external cardiac pacemakers with		
IN nº 09/13					internal power source	12008-03	IEC 60601-2-31
IN nº 09/13					Medical electrical equipment Part_2-		
IN nº 09/13					31: Particular requirements for basic		
	2013	ABNT NBR IEC 60601-2-31	Y	Y	safety and essential performance of		
					external cardiac pacemakers with		
					internal power source	12011-06	IEC 60601-2-31
1					Medical electrical equipment Part_2-		
					31: Particular requirements for basic		
IN nº 09/13	2013	ABNT NBR IEC 60601-2-31	Y	Y	safety and essential performance of		
					external cardiac pacemakers with		
					internal power source	12011-09	IEC 60601-2-31
					Medical electrical equipment; part_2:		
IN nº 09/13	2001	ABNT NBR IEC 60601-2-32	Y	Y	particular requirements for the safety of		
					X-ray equipment	1994-03	IEC 60601-2-32
					Medical electrical equipment Part_2-33:		
					Particular requirements for the basic safety		
					and essential performance of magnetic		
					resonance equipment for medical diagnosis	2010-03	IEC 60601-2-33
					Medical electrical equipment Part_2-		
					33: Particular requirements for the basic		
					safety and essential performance of		
					magnetic resonance equipment for		
					medical diagnosis	2012-03	IEC 60601-2-33
					Medical electrical equipment Part_2-		
IN nº 09/13			Y	Y			
		There is no national			invasive blood pressure monitoring		
		reference			equipment	42011-05	IEC 60601-2-34
					Medical electrical equipment Part_2:		
IN nº 09/13	2006		V	V	Particular requirements for the safety of		
IN Nº 09/13	2006	ABINT INDR IEC 60601-2-36	ř	ř	equipment for extracorporeally induced		
					lithotripsy	1997-03	IEC 60601-2-36
					Medical electrical equipment Part_2-		
IN nº 09/13			Y	Y			
		There is no national					
		reference				2007-08	IEC 60601-2-37
11	2006	reference ABNT NBR IEC 60601-2-36 There is no national	Y	Y	Medical electrical equipment Part_2: Particular requirements for the safety of equipment for extracorporeally induced lithotripsy	1997-03	

IEC 60601-2-3\$2007-11	N	Medical electrical equipment Part_2- 39: Particular requirements for basic safety and essential performance of peritoneal dialysis equipment	Y	Y	ABNT NBR IEC 60601-2-39	2010	IN nº 09/13
IEC 60601-2-4 2010-12	N	Medical electrical equipment Part_2- 4: Particular requirements for basic safety and essential performance of cardiac defibrillators	Υ	Y	There is no national reference		IN nº 09/13
IEC 60601-2-4(1998-02	N	Medical electrical equipment Part_2- 40: Particular requirements for the safety of electromyographs and evoked response equipment	Y	Y	ABNT NBR IEC 60601-2-40	1998	IN nº 09/13
IEC 60601-2-412009-08	N	Medical electrical equipment Part_2- 41: Particular requirements for basic safety and essential performance of surgical luminaires and luminaires for diagnosis	Y	Y	ABNT NBR IEC 60601-2-41	2012	IN nº 09/13
IEC 60601-2-432010-03	N	Medical electrical equipment Part_2- 43: Particular requirements for basic safety and essential performance of X- ray equipment for interventional procedures	Y	Y	ABNT NBR IEC 60601-2-43	2012	IN nº 09/13
IEC 60601-2-442009-02	N	Medical electrical equipment Part_2- 44: Particular requirements for the basic safety and essential performance of X- ray equipment for computed tomography	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-442010-05	N	Medical electrical equipment Part_2- 44: Particular requirements for the basic safety and essential performance of X- ray equipment for computed tomography	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-452011-02	N	Medical electrical equipment Part_2- 45: Particular requirements for the basic safety and essential performance of mammographic X-ray equipment and mammographic stereotactic devices	Y	Y	ABNT NBR IEC 60601-2-45	2013	IN nº 09/13

IEC 60601-2-462	2010-12	Ν	Medical electrical equipment Part_2- 46: Particular requirements for the basic safety and essential performance of operating tables	Y	Y	ABNT NBR IEC 60601-2-46	2012	IN nº 09/13
IEC 60601-2-472	2012-02	N	Medical electrical equipment Part_2- 47: Particular requirements for the basic safety and essential performance of ambulatory electrocardiographic systems	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-4§2		N	Medical electrical equipment Part_2- 49: Particular requirements for the basic safety and essential performance of multifunction patient monitoring equipment	Y	Y	There is no national reference		IN nº 09/13
IEC 60601-2-5 2	2009-07	N	Medical electrical equipment Part_2- 5: Particular requirements for basic safety and essential performance of ultrasonic physiotherapy equipment	Y	Y	ABNT NBR IEC 60601-2-5	2012	IN nº 09/13
IEC 60601-2-5(2	2009-03	N	Medical electrical equipment Part_2- 50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment	Y	Y	ABNT NBR IEC 60601-2-50	2010	IN nº 09/13
IEC 60601-2-5(2	2010-08	N	Medical electrical equipment Part_2- 50: Particular requirements for the basic safety and essential performance of infant phototherapy equipment	Y	Y	ABNT NBR IEC 60601-2-50	2010	IN nº 09/13
IEC 60601-2-522	2009-12	N	Medical electrical equipment Part_2- 52: Particular requirements for the basic safety and essential performance of medical beds	Y	Y	ABNT NBR IEC 60601-2-52	2013	IN nº 09/13
IEC 60601-2-522		N	Medical electrical equipment Part_2- 52: Particular requirements for the basic safety and essential performance of medical beds	Y	Y	ABNT NBR IEC 60601-2-52	2013	IN nº 09/13
IEC 60601-2-522		N	Medical electrical equipment Part_2- 52: Particular requirements for the basic safety and essential performance of medical beds; Technical Corrigendum_1	Y	Y	ABNT NBR IEC 60601-2-52	2013	IN nº 09/13

IEC 60601-2-542009-06	N	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 radioscopy	Y	Y	ABNT NBR IEC 60601-2-54	2011	IN nº 09/13
IEC 60601-2-542010-03	Ν	Medical electrical equipment Part_2- 54: Particular requirements for the basic safety and essential performance of X- ray equipment for radiography and radioscopy	Y	Y	ABNT NBR IEC 60601-2-54	2011	IN nº 09/13
IEC 60601-2-542011-06	N	Medical electrical equipment Part_2- 54: Particular requirements for the basic safety and essential performance of X- ray equipment for radiography and radioscopy	Y	Y	ABNT NBR IEC 60601-2-54	2011	IN nº 09/13
IEC 60601-2-572011-01	N	Medical electrical equipment Part_2- 57: Particular requirements for the basic safety and essential performance of non- laser light source equipment intended for therapeutic, diagnostic, monitoring and cosmetic/aesthetic use					
IEC 60601-2-6 1984	N	Medical electrical equipment. Part 2: Particular requirements for the safety of microwave therapy equipment	Ν	N			
IEC 60601-2-7 1998-02	N	Medical electrical equipment Part_2- 7: Particular requirements for the safety of high-voltage generators of diagnostic X-ray generators	Y	Y	ABNT NBR IEC 60601-2-7	2001	IN nº 09/13
IEC 60601-2-8 2010-11	N	Medical electrical equipment Part 2-8: Particular requirements for the basic safety and essential performance of therapeutic X-ray equipment operating in the range 10_kV to 1_MV					
IEC 60601-2-8 1997-08	N	Medical electrical equipment Part_2: Particular requirements for the safety of therapeutic X-ray equipment in the range 10_kV to 1_MV; Amendment_1					

					[1
			Medical electrical equipment Part_2-			
			8: Particular requirements for the safety			
			of therapeutic X-ray equipment			
IEC 60601-2-8	1999-04	N	operating in 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	N	equipment			
			Electrical and loading characteristics of			
			X-ray tube assemblies for medical			
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 00027	2001-00		Electroacoustics Audiometric			
			equipment Part_1: Equipment for pure-			
IEC 60645-1	2012-02	Ν	tone audiometry			
IEC 00043-1	2012-02	IN				
	1000 11		Audiometers; part_2: equipment for			
IEC 60645-2	1993-11	N	speech audiometry			
			Electroacoustics Audiometric			
			equipment Part_3: Test signals of			
IEC 60645-3	2007-03	N	short 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	Ν	responses			
	-		Medical electrical equipment			
			Characteristics and test conditions of			
			radionuclide imaging devices Anger			
IEC 60789	2005-10	Ν	type gamma cameras			
	2000-10	IN	type gamina cameras			 I

			Medical electrical equipment				
			Characteristics and test conditions of				
			radionuclide imaging devices Anger				
IEC 60789 Co	ri 2009-10	N	type gamma cameras; Corrigendum_1				
			Determination of the maximum				
			symmetrical radiation field from a				
			rotating anode X-ray tube for medical				
IEC 60806	1984	Ν	diagnosis				
			Medical electrical equipment Medical				
			electron accelerators Functional				
IEC 60976	2007-10	N	performance characteristics				
120 00070	2007 10		Safety requirements for electrical				
			equipment for measurement, control				
			and laboratory use Part_2-040:				
			Particular requirements for sterilizers				
			and washer-disinfectors used to treat				
IEC 61010-2-0	42005-04	Ν	medical materials				
			Safety requirements for electrical				
			equipment for measurement, control				
			and laboratory use Part_2-101:	Y	Y		IN nº 09/13
			Particular requirements for in vitro			There is no national	
IEC 61010-2-1	2002-01	N	diagnostic (IVD) medical equipment			reference	
			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 Co	ri 2008-08	N	ultrasonic equipment; Corrigendum_1				
	2000 00		Radiotherapy simulators; functional				
IEC 61168	1993-12	Ν	performance characteristics				
			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				

			Evoluction and routing testing in	
			Evaluation and routine testing in	
			medical imaging departments Part_2-	
			6: Constancy tests Imaging	
	0000 44		performance of computed tomography	
IEC 61223-2-6	2006-11	N	X-ray equipment	
			Evaluation and routine testing in	
			medical imaging departments Part_3-	
			2: Acceptance tests Imaging	
			performance of mammographic X-ray	
IEC 61223-3-2	2007-07	N	equipment	
			Evaluation and routine testing in medical	
			imaging departments Part_3-4:	
IEC 61223-3-4	2000-03	N	Acceptance tests Imaging performance of dental X-ray equipment	
IEC 01223-3-4	2000-03	IN	Evaluation and routine testing in	
			medical imaging departments Part_3-	
			5: Acceptance tests Imaging	
			performance of computed tomography	
IEC 61223-3-5	2004.02	N	X-ray equipment	
IEC 01223-3-5	2004-06	N		
			Evolution and routing togting in	
			Evaluation and routine testing in	
			medical imaging departments Part_3-	
			5: Acceptance tests Imaging performance of computed tomography	
IEC 61223-3-5	2006.02	N		
IEC 01223-3-5	2006-03	N	X-ray equipment; Corrigendum_1	
IEC 61252 Edit	+ 2002 02	N	Electroacoustics Specifications for personal sound exposure meters	
IEC 01252 EUI	12002-03	N	personal sound exposure meters	
			Madical alastrical aquinment	
			Medical electrical equipment	
			Characteristics of electro-optical X-ray	
150 04000 4	1001.07	N	image intensifiers Part_1:	
IEC 61262-1	1994-07	N	Determination of the entrance field size	
			Medical electrical equipment	
	1		Medical electrical equipment	
			Characteristics of electro-optical X-ray	
	4004.07		image intensifiers Part_2:	
IEC 61262-2	1994-07	N	Determination of the conversion factor	

						1	
			Medical electrical equipment				
			Characteristics of electro-optical X-ray				
			image intensifiers Part_3:				
			Determination of the luminance				
			distribution and luminance non-				
IEC 61262-3	1994-07	N	uniformity				
			Medical electrical equipment				
			Characteristics of electro-optical X-ray				
			image intensifiers Part_4:				
IEC 61262-4	1994-07	Ν	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				
			image intensifiers Part-7:				
			Determination of the modulation transfer				
IEC 61262-7	1995-09	Ν	function				
			Ultrasonics Hand-held probe Doppler				
			foetal heartbeat detectors				
			Performance requirements and				
IEC 61266	1994-12	Ν	methods of 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				
	1334-03	IN	methous for describing performance		Į	<u> </u>	Į

IEC 61326-2-6	2005-12	Ν	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	2007-09	Ν	Electrical equipment for measurement, control and laboratory use, control and laboratory use EMC requirements Part_2-6: Particular requirements In- vitro diagnostic (IVD) medical equipment; Corrigendum_1			
IEC 61331-1	1994-10	N	Protective devices against diagnostic medical X-radiation Part_1: Determination of attenuation properties of materials			
IEC 61331-2	1994-10	N	Protective devices against diagnostic medical X-radiation Part_2: Protective glass plates			
IEC 61331-3	1998-11	N	Protective devices against diagnostic medical X-radiation Part_3: Protective clothing and protective devices for gonads			
IEC 61391-1	2006-07	N	Ultrasonics Pulse echo scanners Part_1: Techniques for calibrating spatial measurement systems and measurement of system point-spread function response			
			Ultrasonics Pulse-echo scanners Part_2: Measurement of maximum depth of penetration and local dynamic			
IEC 61391-2	2010-01	<u> </u>	range Electroacoustics Equipment for the measurement of real-ear acoustical characteristics of hearing aids			
IEC 61674 AM	[2002-06	N	Medical electrical equipment Dosimeters with ionization chambers and/or semi-conductor detectors as used in X-ray diagnostic imaging; Amendment_1			

			Radionuclide imaging devices			
			Characteristics and test conditions			
IEC 61675-1	1998-02	N	Part_1: Positron emission tomographs			
			Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_1: Positron emission tomographs;			
IEC 61675-1 A	2008-04	N	Amendment_1			
			Dedienvelide imperient devices			
			Radionuclide imaging devices Characteristics and test conditions			
IEC 61675-1 E		N	Part_1: Positron emission tomographs			
	12008-06	IN	Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_2: Single photon emission			
IEC 61675-2	1998-01	Ν	computed tomographs			
120 010/02	1000 01					
			Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_2: Single photon emission			
IEC 61675-2 A	2004-12	N	computed tomographs; Amendment_1			
			Radionuclide imaging devices			
			Characteristics and test conditions			
			Part_2: Single photon emission			
IEC 61675-2 E	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-			
	0000.00		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 AM	12008-11	N	Amendment 1			
		IN			1	

			Medical electrical equipment			
			Dosimetric instruments used for non-			
			invasive measurement of X-ray tube			
IEC 61676 Edit	2009-01	Ν	voltage in diagnostic radiology			
			Medical electrical equipment Dosimetric			
			instruments used for non-invasive			
			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	Ν	0,5_MHz to 5_MHz			
			Ultrasonics Pressure pulse			
IEC 61846	1998-04	Ν	lithotripters Characteristics of fields			
			Ultrasonics Surgical systems			
			Measurement and declaration of the			
IEC 61847	1998-01	Ν	basic output characteristics			
			Medical electrical equipment			
			Requirements for the safety of			
			radiotherapy treatment planning			
IEC 62083	2009-09	N	systems			
			Medical electrical equipment			
			Characteristics of digital X-ray imaging			
			devices Part_1: Determination of the			
IEC 62127.1	2003-10	N	detective quantum efficiency			-
			Medical electrical equipment			
			Characteristics of digital X-ray imaging			
	0000 40		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			
IEC 62220-1-2	2007-06	N	Detectors used in mammography			
			Medical electrical equipment			
			Characteristics of digital X-ray imaging			
			devices Part_1-3: Determination of			
			the detective quantum efficiency			
IEC 62220-1-3	2008-06	N	Detectors used in dynamic imaging			

	1	1				
			Medical electrical equipment Safety of			
	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			
IEC 62353	2007-05	N	medical electrical equipment			
			Little and the Etablish are starting to a			
			Ultrasonics Field characterization			
			Test methods for the determination of			
			thermal and mechanical indices related			
IEC 62359	2010-10	N	to medical diagnostic ultrasonic fields			
			Litragonica Field characterization			
			Ultrasonics Field characterization			
			Test methods for the determination of			
			thermal and mechanical indices related			
IEC 62359 Co	12011-03	N	to medical diagnostic ultrasonic fields			
			Medical devices Application of			
IEC 62366	2007-10	N	usability engineering to medical devices			
IEC 02300	2007-10	IN	Magnetic resonance equipment for medical			
			imaging Part_1: Determination of essential			
IEC 62464-1	2007-01	N	image quality parameters			
	2007 01		Magnetic resonance equipment for medical			
			imaging Part_2: Classification criteria for			
IEC 62464-2	2010-11	N	pulse sequences			
			Electroacoustics Audio-frequency			
			induction loop systems for assisted hearing			
			Part_1: Methods of measuring and			
			specifying the performance of system			
IEC 62489-1	2010-01	N	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 loop for assessing			
			conformity with guidelines on limits for			
IEC 62489-2	2011-01	N	human exposure			

			Medical electrical equipment					
			Exposure index of digital X-ray imaging					
			systems Part_1: Definition and					
IEC 62494-1	2008-08	N	requirements of 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					
IEC 80001-1	2010-10	N	and activities					
			Medical electrical equipment Part_2-					
			30: Particular requirements for basic					
			safety and essential performance of	Y	Y			IN nº 09/13
			automated non-invasive			There is no national		
IEC 80601-2-3	2009-01	N	sphygnomanometers			reference		
			Medical electrical equipment Part_2-					
			30: Particular requirements for basic	Y	Y			IN nº 09/13
			safety and essential performance of	I	T			IN 11º 09/13
			automated non-invasive			There is no national		
IEC 80601-2-3	2010-01	N	sphygnomanometers; Corrigendum_1			reference		
			Medical electrical equipment Part_2-					
			35: Particular requirements for the basic					
			safety and essential performance of	Y	Y	ABNT NBR IEC 80601-2-35	2013	IN nº 09/13
			heating devices using blankets, pads	I	1	ABINT NBICIEC 00001-2-33	2015	1111 03/13
			and mattresses and intended for					
IEC 80601-2-3	2009-10	N	heating in medical use					
			Medical electrical equipment Part_2-					
			35: Particular requirements for the basic					
			safety and essential performance of	Y	Y	ABNT NBR IEC 80601-2-35	2013	IN nº 09/13
			heating devices using blankets, pads	·	•		2010	
			and mattresses and intended for					
IEC 80601-2-3	2012-03	N	heating in medical use					
			Medical electrical equipment Part_2-					
			58: Particular requirements for basic			ABNT NBR ISO/IEC 80601-2-		
			safety and essential performance of	Y	Y	58	2013	IN nº 09/13
			lens removal devices and vitrectomy					
IEC 80601-2-5	ą2008-10	N	devices for ophthalmic surgery					

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			Medical electrical equipment Part_2-	
			59: Particular requirements for basic	
			safety and essential performance of	
			screening thermographs for human	
IEC 80601-2-59	2008-10	N	febrile temperature screening	
			Medical electrical equipment Part_2-	
			59: Particular requirements for basic	
			safety and essential performance of	
			screening thermographs for human	
			febrile temperature screening;	
IEC 80601-2-59	2009-04	N	Corrigendum_1	
			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 for the safe use of laser beams	
IEC/TR 60825-8	2006-12	N	on humans	
			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	
			Cuidelines for administrative medical	
			Guidelines for administrative, medical,	
			and nursing staff concerned with the	
	2002 02	K 1	safe use of medical electrical equipment	
IEC/TR 60930	2008-09	N	and medical electrical systems	
			Medical electrical equipment Medical	
			electron accelerators Guidelines for	
IEC/TR 60977	2008-07	N	functional performance characteristics	
		1	Guidelines for the development and use	
			of medical electrical equipment	
IEC/TR 61258	2008-08	N	educational materials	
			High frequency surgical equipment	
IEC/TR 61289	2011-11	N	Operation and maintenance	
0/ 110 01200				

			Nuclear medicine instrumentation				
			Routine tests Part_2: Scintillation				
			cameras and single photon emission				
IEC/TR 61948-	2001-02	N	computed tomography imaging				
1EC/TK 01940-	2001-02	IN	Nuclear medicine instrumentation				
			Routine tests Part_3: Positron				
IEC/TR 61948-	2005 07	N					
IEC/TR 01948-	2005-07	N	emission tomographs				
			Nuclear medicine instrumentation				
	0000 44	N	Routine tests Part_4: Radionuclide				
IEC/TR 61948-	2006-11	N	calibrators			-	
			Medical electrical equipment				
			Guidelines for implementation of				
IEC/TR 62266	2002-03	N	DICOM in radiotherapy				
			Considerations of unaddressed safety				
			aspects in the second edition of				
			IEC_60601-1 and proposals for new				
IEC/TR 62296	2009-01	N	requirements				
			Mapping between the clauses of the				
			third edition of IEC_60601-1 and the				
IEC/TR 62348	2006-05	N	1988 edition as amended				
			General testing procedures for medical				
IEC/TR 62354	2009-10	N	electrical equipment				
			Requirements for measurement				
			standards for high intensity therapeutic				
IEC/TR 62649	2010-04	Ν	ultrasound (HITU) devices				
			Audio, video and multimedia systems and				
			equipment Activities and considerations				
IEC/TR 62678	2010-10	N	related to accessibility and usability				
			Medical device software Part_1: Guidance				
IEC/TR 80002-1	2000.00	Ν	on the application of ISO_14971 to medical device software				
ILC/TR 00002-1	2009-09	IN					
			Radiotherapy simulators; guidelines for				
IEC/TR2 61170	1002 12	N	functional performance characteristics				
	1993-12	IN	Evaluation and routine testing in				
IEC/TR2 61223	1002.07	N	medical imaging departments; part_1:				
1EC/182 01223	1993-07	N	general aspects				
			Ultrasonics Real-time pulse-echo				
			systems Test procedures to				
	4000 07	N	determine the performance				
IEC/TR2 61390	1996-07	N	specifications		l		

			Fundamental aspects of safety				
			standards for medical electrical				
IEC/TR3 60513	1994-01	N	equipment				
	1004 01	11					
			Cardiac defibrillators; cardiac				
IEC/TR3 61288	1993-10	N	defibrillators-monitors; part_1: operation				
120/11001200			Cardiac defibrillators; cardiac				
			defibrillators-monitors; part_2:				
IEC/TR3 61288	1993-10	N	maintenance				
			Medical electrical equipment Digital				
			imaging and communications in				
			medicine (DICOM) Radiotherapy				
IEC/TR3 61852	1998-04	N	objects				
120/1100/01002	1000 04	11	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				
	0005.00		removable dentures Part_1: Materials for				
ISO 10139-1	2005-02	N	short-term use Dentistry Soft lining materials for				
			removable dentures Part_1: Materials for				
ISO 10139-1 Teo	2006-03	N	short-term use; Technical Corrigendum_1				
			Dentistry Soft lining materials for				
			removable dentures Part_2: Materials for				
ISO 10139-2	2009-08	N	long-term use				
			Health informatics Messages and				
100 10150			communication Web access reference				
ISO 10159	2011-12	N	manifest Dentistry Corrosion test methods for				
ISO 10271	2011-08	N	metallic materials				
100 10271	2011/00		Single-use sterile rubber surgical				
ISO 10282	2002-09	N	gloves Specification	Y	Y		RDC 55/2011
100 10202	2002-03	IN	gioves opecilication		L	ļ	 1

			Single-use sterile rubber surgical				
l			gloves Specification; Technical	Y	Y		RDC 55/2011
ISO 10282 Tec	2005 06	Ν	Corrigendum_1	•			1100 00/2011
150 10202 160	2003-00	IN	Ophthalmic optics Semi-finished				
1			spectacle lens blanks Part_1:				
1							
			Specifications for single-vision and				
ISO 10322-1	2006-02	N	multifocal lens blanks				
l			Ophthalmic optics Semi-finished				
l			spectacle lens blanks Part_2:				
			Specifications for progressive power				
ISO 10322-2	2006-02	N	lens blanks				
l			Dental rotary instruments; bore diameters				
ISO 10323	1991-11	N	for discs and wheels				
l			Prosthetics Structural testing of lower-limb				
100 40000	0000 40	N	prostheses Requirements and test methods				
ISO 10328	2006-10	N					
l			Implants for surgery Malleable wires				
	4004.00		for use as sutures and other surgical				
ISO 10334	1994-08	N	applications				
			Ophthalmic instruments Refractor				
ISO 10341	2009-07	N	heads				
l			Ophthalmic instruments Eye				
ISO 10342	2010-06	N	refractometers				
l			Ophthalmic instruments				
ISO 10343	2009-07	N	Ophthalmometers				
l			Dentistry Contents of technical file for				
ISO 10451	2010-06	N	dental implant systems				
			Dentistry Polymer-based crown and bridge				
ISO 10477	2004-10	N	materials				
l			Pressure regulators for use with				
l			medical gases Part_1: Pressure				
l			regulators and pressure regulators with				
ISO 10524-1	2006-02	N	flow-metering devices				
l			Pressure regulators for use with				
l			medical gases Part_2: Manifold and				
ISO 10524-2	2005-05	Ν	line pressure regulators				
			Pressure regulators for use with				
l			medical gases Part_3: Pressure				
			regulators integrated with cylinder				
ISO 10524-3	2005-05	Ν	valves				

			Pressure regulators for use with	T		
			medical gases Part_4: Low-pressure			
100 40504 4	2000.00	N	S			
ISO 10524-4	2008-06	N	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:			
100 105 10 1	0004.07		Requirements and test methods for all			
ISO 10542-1	2001-07	N	systems			
			Technical systems and side for dischlod or			
			Technical systems and aids for disabled or handicapped persons Wheelchair tiedown			
			and occupant-restraint systems Part_2:			
ISO 10542-2	2001-07	N	Four-point strap-type tiedown systems			
100 10042 2	2001 01		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			
			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			
			and occupant-restraint systems Part_5:			
ISO 10542-5	2004-04	N	Systems for specific wheelchairs			
			Sterile, single-use intravascular			
			catheters Part_1: General			
ISO 10555-1	1995-06	N	requirements			
			Sterile, single-use intravascular			
			catheters Part_1: General			
ISO 10555-1 A	AI 1999-07	N	requirements; Amendment_1			
			Sterile, single-use intravascular			
			catheters Part_1: General			
ISO 10555-1 A	2004-05	N	requirements; Amendment_2			
32 3000 17			Sterile, single-use intravascular			
			catheters Part_2: Angiographic			
ISO 10555-2	1996-06	N	catheters			
100 10000-2	1330-00	IN	Sterile, single-use intravascular			
100 40555 0 7			catheters Part_2: Angiographic			
ISO 10555-2 1	102002-06	N	catheters; Technical Corrigendum_1			

			Sterile, single-use intravascular					
			catheters Part_3: Central venous					
ISO 10555-3	1996-06	N	catheters					
100 10000-0	1330-00		Sterile, single-use intravascular					
			catheters Part_3: Central venous					
ISO 10555-3 T	2002.06	N	catheters; Technical Corrigendum_1					
130 10555-5 1	62002-00	IN	Sterile, single-use intravascular					
	1000.00	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 T	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 A	1999-01	N	peripheral catheters; Amendment_1					
			Sterile, single-use intravascular					
			catheters Part_5: Over-needle					
			peripheral catheters; Technical					
ISO 10555-5 T	2002-06	N	Corrigendum_1					
			Dental equipment High- and medium-					
ISO 10637	1999-08	N	volume suction systems					
			Dentistry Powered polymerization					
100 40050 4			activators Part_1: Quartz tungsten					
ISO 10650-1	2004-11	N	halogen lamps Dentistry Powered polymerization					
			activators Part_2: Light-emitting diode					
ISO 10650-2	2007-09	Ν	(LED) lamps					
130 10030-2	2007-03		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						
150 10651-2	2004-07	IN	dependent patients					
			Lung ventilators for medical use					
100 40054 0	1007.01	K I	Part_3: Particular requirements for					
ISO 10651-3	1997-01	N	emergency and transport ventilators					
			Lung ventilators Part_4: Particular	N/	N/			IN - 0 00/4 0
			requirements for operator-powered	Y	Y	ABNT NBR ISO 10651-4		IN nº 09/13
ISO 10651-4	2002-03	N	resuscitators				2011	

	1					1
			Lung ventilators for medical use			
			Particular requirements for basic safety			
			and essential performance Part_5:			
100 10051 5	2006-02	N				
ISO 10651-5	2006-02	N	Gas-powered emergency resuscitators			
			Lung ventilators for medical use			
			Particular requirements for basic safety			
			and essential performance Part_6:			
ISO 10651-6	2004-07	N	Home-care ventilatory support devices			
130 10031-0	2004-07	IN	Ophthalmic optics Spectacle frames			
			and sunglasses electronic catalogue			
			and identification Part_1: Product			
			identification and electronic catalogue			
100 10005 1	2011-12	N	product hierarchy			
ISO 10685-1 ISO 10873	2010-09	N N	Dentistry Denture adhesives			
130 10073	2010-03		Optics and optical instruments			
			Operation microscopes Part_1:			
ISO 10936-1	2000-06	N	Requirements and test methods			
100 10000 1	2000 00		Optics and photonics Operation			
			microscopes Part_2: Light hazard			
			from operation microscopes used in			
ISO 10936-2	2010-01	N	ocular surgery			
100 10000 2	2010 01		Ophthalmic instruments Chart			
ISO 10938	1998-05	N	projectors			
			Ophthalmic instruments Slit-lamp			
ISO 10939	2007-02	N	microscopes			
			Ophthalmic instruments Fundus			
ISO 10940	2009-08	N	cameras			
			Ophthalmic instruments Direct			
ISO 10942	2006-06	Ν	ophthalmoscopes			
			Ophthalmic instruments Indirect			
ISO 10943	2011-08	Ν	ophthalmoscopes			
			Ophthalmic instruments			
ISO 10944	2009-08	N	Synoptophores			
			Caps made of aluminium-plastics			1
			combinations for infusion bottles and			
			injection vials Requirements and test			
ISO 10985	2009-02	Ν	methods			
					•	

			Biological evaluation of medical					
			devices Part_1: Evaluation and		_		0010	
			testing within a risk management	Y	Р	ABNT NBR ISO	2013	RDC 16/2012
ISO 10993-1	2009-10	Ν	process					
			Biological evaluation of medical					
			devices Part_1: Evaluation and					
			testing within a risk management					
ISO 10993-1 T	2010-06	N	process; Technical Corrigendum_1					
			Biological evaluation of medical					
			devices Part_10: Tests for irritation					
ISO 10993-10	2010-08	N	and skin sensitization					
	2010 00		Biological evaluation of medical					
			devices Part_11: Tests for systemic					
ISO 10993-11	2006-08	N	toxicity					
100 10000 11	2000 00	11	Biological evaluation of medical					
			devices Part_12: Sample preparation					
ISO 10993-12	2007-11	N	and reference materials					
100 10333-12	2007-11	11	Biological evaluation of medical					
			devices Part_13: Identification and					
			quantification of degradation products					
ISO 10993-13	2010-06	Ν	from polymeric medical devices					
100 10333-13	2010-00	11	Biological evaluation of medical					
			devices Part_14: Identification and					
			quantification of degradation products					
ISO 10993-14	2001-11	Ν	from ceramics					
100 10333-14	2001-11	11	Biological evaluation of medical					
			devices Part_15: Identification and					
			quantification of degradation products					
ISO 10993-15	2000-12	N	from metals and alloys					
100 10333-13	2000-12	11	Biological evaluation of medical					
			devices Part_16: Toxicokinetic study					
			design for degradation products and					
ISO 10993-16	2010-02	N	leachables					
130 10993-10	2010-02	11	Biological evaluation of medical					
			devices Part_17: Establishment of					
			allowable limits for leachable					
ISO 10993-17	2002-12	N	substances					
100 10000-17	2002-12	11	Biological evaluation of medical					
			devices Part_18: Chemical					
ISO 10993-18	2005-07	Ν	characterization of materials					
100 10993-18	2003-07	IN						

			Biological evaluation of medical	
			devices Part_2: Animal welfare	
ISO 10993-2	2006-07	Ν	requirements	
100 10990-2	2000-07	11		
			Biological evaluation of medical	
			devices Part_3: Tests for genotoxicity,	
ISO 10993-3	2003-10	Ν	carcinogenicity and reproductive toxicity	
100 10330-0	2003-10	11	Biological evaluation of medical	
			devices Part_4: Selection of test for	
ISO 10993-4	2002-10	Ν	interactions with blood	
100 10333-4	2002-10	11	Biological evaluation of medical	
			devices Part_4: Selection of tests for	
ISO 10993-4 A	2006 07	N	interactions with blood	
130 10993-47	12000-07	IN	Biological evaluation of medical	
			devices Part_5: Tests for in vitro	
ISO 10993-5	2009-06	N	cytotoxicity	
130 10993-5	2009-00	IN	Biological evaluation of medical	
			devices Part_6: Tests for local effects	
ISO 10993-6	2007-04	N	after implantation	
130 10993-0	2007-04	IN	Biological evaluation of medical	
			devices Part_7: Ethylene oxide	
ISO 10993-7	2008-10	N	sterilization residuals	
130 10993-7	2000-10	IN	Biological evaluation of medical	
			devices Part_7: Ethylene oxide	
			sterilization residuals; Technical	
ISO 10993-7 1	2000 11	N	Corrigendum_1	
130 10993-7 1	2009-11	IN	Biological evaluation of medical	
			devices Part_9: Framework for	
			identification and quantification of	
ISO 10993-9	2009-12	N	potential degradation products	
150 10993-9	2003-12		Prefilled syringes; part_1: glass	
			cylinders for dental local anaesthetic	
ISO 11040-1	1992-11	N	cartridges	
130 11040-1	1992-11	IN	Prefilled syringes Part_2: Plunger	
			stoppers for dental local anaesthetic	
ISO 11040-2	2011-04	N	cartridges	
130 11040-2	2011-04	IN	Prefilled syringes Part_3: Seals for dental	
ISO 11040-3	2012-01	N	local anaesthetic cartridges	
			Prefilled syringes Part_4: Glass	
ISO 11040-4	2007-02	Ν	barrels for injectables	

			Prefilled syringes Part_5: Plunger			
ISO 11040-5	2012-01	N	stoppers for injectables			
150 11040-5	2012-01	IN	Sterile single-use intravascular catheter			
100 11070	1000.05	N	5			
ISO 11070	1998-05	N	introducers Health informatics Point-of-care medical			
ISO 11073-9010	2008 01	N	device communication Part_90101: Analytical instruments Point-of-care test			
130 1107 3-9010	2008-01	IN	Health informatics Standard			
			communication protocol Part_91064:			
ISO 11073-9106	2009-05	Ν	Computer-assisted electrocardiography			
			Sterilization of health care products			
			Ethylene oxide Part_1: Requirements			
			for development, validation and routine			
			control of a sterilization process for			
100 44405 4	0007.05	N				
ISO 11135-1	2007-05	N	medical devices			
			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	N	medical devices			
			Sterilization of health care products			
			Radiation Part_2: Establishing the			
ISO 11137-2	2012-03	N	sterilization dose			
			Sterilization of health care products			
			Radiation Part_3: Guidance on			
ISO 11137-3	2006-04	N	dosimetric aspects			
			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			
	2000 01		Sterilization of health care products			
			Biological indicators Part_3: Biological			
			indicators for moist heat sterilization			
ISO 11138-3	2006-07	Ν	processes			
100 11100-0	2000-07	IN	Sterilization of health care products			
			Biological indicators Part_4: Biological			
100 44400 4	0000 07	N 1	indicators for dry heat sterilization			
ISO 11138-4	2006-07	N	processes			

			Sterilization of health care products					
			Biological indicators Part_5: Biological					
			indicators for low-temperature steam					
			and formaldehyde sterilization					
ISO 11138-5	2006-07	N	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					
			Starilization of boolth care products					
			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 T	L2007-11	N	Technical Corrigendum_1					
130 11140-3 1	102007-11	IN	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					
ISO 11140-4	2007-03	Ν	steam penetration					
	2007 00		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	N	Dentistry Amalgam separators					
			Dental equipment Connections for supply					
ISO 11144	1995-05	N	and waste lines					
100 44450	0044.07		Packaging Accessible design General					
ISO 11156	2011-07	N	requirements					
			Single-use medical examination gloves_					
			Part_1: Specification for gloves made	Y	Y			
ISO 11193-1	2008-09	Ν	from rubber latex or rubber solution			ABNT NBR ISO	2009	RDC 55/2011
			Single-use medical examination gloves_					
			Part_2: Specification for gloves made					
ISO 11193-2	2006-11	N	from poly(vinyl chloride)					
			Gas mixers for medical use Stand-	Y	Y	ABNT NBR ISO 11195		IN nº 09/13
ISO 11195	1995-10	N	alone gas mixers	1	•		2000	

ISO 11197	2004-12	N	Medical supply units					
			Walking aids manipulated by both arms					
			Requirements and test methods Part_1:					
ISO 11199-1	1999-08	N	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					
			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					
100 11010	2002 00	11	Assistive products for walking manipulated					
			by one arm Requirements and test					
ISO 11334-1	2007-02	N	methods Part_1: Elbow crutches					
	2001 02		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					
			Transfusion equipment for medical use;					
ISO 1135-3	1986-11	N	Part 3 : Blood-taking set					
150 1155-5	1900-11	IN	Fait 5. Diood-taking Set					
				V	Ň			DDC 04/0044
100 110- 1			Transfusion equipment for medical use_	Y	Y			RDC 04/2011
ISO 1135-4	2012-03	N	Part_4: Transfusion sets for single use					
			Optics and optical instruments	Y	NO	Revision Project ABNT NBR	2013	
ISO 11380	1994-10	N	Ophthalmic optics Formers	•		ISO	2010	
			Optics and optical instruments	Y	NO	ABNT NBR ISO	2003	
ISO 11381	1994-12	N	Ophthalmic optics Screw threads	I	NO	ABINT INDR 130	2003	
			Containers and accessories for					
			pharmaceutical preparations Part_1: Drop-	Y	NO	ABNT NBR ISO	2003	
ISO 11418-1	2005-02	Ν	dispensing glass bottles					
			Containers and accessories for					
			pharmaceutical preparations Part_2:	Y	NO	ABNT NBR ISO	2003	
ISO 11418-2	2005-02	Ν	Screw-neck glass bottles for syrups					
			Containers and accessories for					
			pharmaceutical preparations Part_3:					
			Screw-neck glass bottles (veral) for solid					
ISO 11418-3	2005-02	N	and liquid dosage forms					
			Containers and accessories for					
			pharmaceutical preparations Part_4:					
ISO 11418-4	2005-02	N	Tablet glass bottles					

			Containers and accessories for					
			pharmaceutical preparations Part_5:					
ISO 11418-5	1997-12	N	Dropper assemblies					
			Containers and accessories for					
			pharmaceutical preparations Part_7:					
			Screw-neck vials made of glass tubing for					
ISO 11418-7	1998-10	N	liquid 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:	Y	NO	ABNT NBR ISO	2009	
				I	NO	ABINT INDIC ISO	2009	
	2000 04	N	Requirements for materials, sterile					
ISO 11607-1	2006-04	N	barrier systems and packaging systems					
			Packaging for terminally sterilized					
			medical devices Part_2: Validation	Y	NO	ABNT NBR ISO	1999	
			requirements for forming, sealing and					
ISO 11607-2	2006-04	N	assembly processes					
			Pen-injectors for medical use Part_1:					
			Pen-injectors; Requirements and test					
ISO 11608-1	2000-12	N	methods					
			Pen-injectors for medical use Part_2:					
			Needles; Requirements and test					
ISO 11608-2	2000-12	Ν	methods					
			Pen-injectors for medical use Part_3:					
			Finished cartridges; Requirements and					
ISO 11608-3	2000-12	Ν	test methods					
			Pen-injectors for medical use Part_4:					
			Requirements and test methods for					
			electronic and electromechanical pen-	Y	NO	ABNT NBR ISO	1997	
ISO 11608-4	2006-03	Ν	injectors					
100 11000 4	2000 00		Dentistry Dentifrices Requirements, test					
ISO 11609	2010-09	Ν	methods and marking					
			Quality of dialysis fluid for					
ISO 11663	2009-04	N	haemodialysis and related therapies					
			Packaging Tactile warnings of danger				1	
ISO 11683	1997-10	Ν	Requirements					
							1	
			Anaesthetic and respiratory equipment					
ISO 11712	2009-05	N	Supralaryngeal airways and connectors					
			oup and figure and officiolo		1		1	

	T	Other Weiner Contract Contract States in the					
2006-04	N						
		Sterilization of medical devices					
		Microbiological methods Part_1:					
		Determination of a population of					
		microorganisms on products; Technical					
Ce2007-05	Ν						
2000 11	N						
2003-11							
		Lesson and lesson valated any invest					
2005-02	N						
		Test method and classification for the					
		laser-resistance of surgical drapes					
		and/or patient-protective covers					
2007-05	N	Part_2: Secondary ignition					
		Acoustics Determination of sound					
		immission from sound sources placed close					
2002-10	N						
1996-11	N						
2010-06	N						
2000-03	Ν	manufacturer					
		Ophthalmic implants Intraocular					
2006-07	N	lenses Part_1: Vocabulary					
	2002-10 1996-11 2010-06 2000-03	2007-05 N 2009-11 N 2005-02 N 2007-05 N 2007-05 N 2002-10 N 1996-11 N 2010-06 N 2000-03 N	Sterilization of medical devices Microbiological methods Part_1: Determination of a population of microorganisms on products; Technical 2007-05 N Corrigendum_1 Sterilization of medical devices Microbiological methods Part_2: Tests of sterility performed in the definition, validation and maintenance 2009-11 N of a sterilization process 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser resistance of surgical drapes and/or patient protective covers 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers 2007-05 N Part_2: Secondary ignition Acoustics Determination of sound immission from sound sources placed close for the ear Part_1: Technique using a microphone in a real ear (MIRE technique)	Microbiological methods Part_1: Determination of a population of 2006-04 N microorganisms on products Microbiological methods Part_1: Determination of a population of microorganisms on products; Technical 2007-05 N Corrigendum_1 Sterilization of medical devices Microbiological methods Part_2: Tests of sterility performed in the definition, validation and maintenance 2009-11 N of a sterilization process 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser resistance of surgical drapes and/or patient protective covers 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers and/or patient-protective covers 2007-05 N Part_2: Secondary ignition Acoustics Determination of sound immission from sound sources placed close for the ear Part_1: Technique using a microphone in a real ear (MIRE technique) 2002-10 N Morthobing aids Part_1: Whole- product testing 2010-06 N Mor	Microbiological methods Part_1: Determination of a population of microorganisms on products Sterilization of medical devices Microbiological methods Part_1: Determination of a population of microorganisms on products; Technical 2007-05 N Corrigendum_1 2007-05 N Sterilization of medical devices Microbiological methods Part_2: Tests of sterility performed in the definition, validation and maintenance 2009-11 N of a sterilization process 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser resistance of surgical drapes and/or patient protective covers 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser-resistance of surgical drapes and/or patient protective covers 2007-05 N Part_2: Secondary ignition Acoustics Determinati	2006-04 N Determination of a population of microorganisms on products 2006-04 N Sterilization of medical devices Microbiological methods Part_1: Determination of a population of microorganisms on products, Technical 2007-05 N Corrigendum_1 2007-05 Sterilization of medical devices Microbiological methods Part_2: Tests of sterilization of medical devices Microbiological methods Part_2: Tests of sterilization process 2009-11 N of a sterilization process 2005-02 N Easers and laser-related equipment Test method and classification for the laser resistance of surgical drapes and/or patient protective covers 2005-02 N Part_1: Primary ignition and penetration 2007-05 N Part_1: Primary ignition and penetration 2007-05 N Part_2: Secondary ignition 2007-06 N microphone in a real ear (MIRE technique) 1996-11 N microphone in a real ear (MIRE technique) 1996-11 N product testing 2010-06 N nealevistruments	2006-04 N 2006-04 N Interocorganisms on products Sterilization of maclical devices Microbiological methods Part_1: Determination of a population of microorganisms on products; Technical 2007-05 N Corrigendum_1 2009-11 N of a sterilization of maclical devices Microbiological methods Part_2: Tests of sterility performed in the definition, validation and maintenance 2009-11 N of a sterilization process 2005-02 N Part_1: Primary ignition and penetration Lasers and laser-related equipment Test method and classification for the laser resistance of surgical drapes and/or patient protective covers and/or patient-protective

			Ophthalmic implants Intraocular					
			lenses Part_10: Phakic intraocular					
ISO 11979-10	2006-08	Ν	lenses					
			Ophthalmic implants Intraocular					
			lenses Part_2: Optical properties and					
ISO 11979-2	1999-12	Ν	test methods					
			Ophthalmic implants Intraocular					
			lenses Part_2: Optical properties and					
ISO 11979-2 T	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	Y	NO	ABNT NBR ISO	2009	
ISO 11979-4	2008-12	N	information					
			Ophthalmic implants Intraocular	Y	NO	ABNT NBR ISO	2009	
ISO 11979-5	2006-06	N	lenses Part_5: Biocompatibility	•			2000	
			Ophthalmic implants Intraocular					
			lenses Part_6: Shelf-life and transport	Y	NO	ABNT NBR ISO	2008	
ISO 11979-6	2007-07	N	stability					
			On hith a locial international lateral avelan	Y	NO	ABNT NBR ISO	1998	
100 44070 7	0000 05	N	Ophthalmic implants Intraocular	ř	NO	ABINT INBR ISO	1998	
ISO 11979-7	2006-05	N	lenses Part_7: Clinical investigations Ophthalmic implants Intraocular					
				Y	NO	ABNT NBR ISO	1998	
ISO 11979-7 A	2012.01	N	lenses Part_7: Clinical investigations; Amendment_1	ř	NO	ABINT INBR ISO	1998	
150 11979-7 A	2012-01	IN	Ophthalmic implants Intraocular					
			lenses Part_8: Fundamental					
ISO 11979-8	2006-07	N	requirements					
100 11979-0	2000-07		Ophthalmic implants Intraocular					
			lenses Part_8: Fundamental					
ISO 11979-8 A	2011-05	N	requirements; Amendment_1					
			Ophthalmic implants Intraocular					
			lenses Part_9: Multifocal intraocular					
ISO 11979-9	2006-09	Ν	lenses					
ISO 11980	2009-10	N	•					
			Ophthalmic optics Contact lenses and contact lens care products Guidance for clinical investigations					

			Ophthalmic optics Contact lenses and					
			contact lens care products					
			Determination of physical compatibility					
			of contact lens care products with					
ISO 11981	2009-07	Ν	contact lenses					
			Ophthalmic optics Contact lenses					
			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					
			Determination of preservative uptake					
ISO 11986	2010-11	Ν	and release					
			Ophthalmic optics Contact lenses	Y	NO			
ISO 11987	1997-12	Ν	Determination of shelf-life	ř	NO			
			Ophthalmic optics Contact lenses					
			Determination of shelf-life; Technical					
ISO 11987 Tec	1998-04	Ν	Corrigendum_1					
			Lasers and laser-related equipment					
			Determination of laser resistance of					
			tracheal tubes Part_1: Tracheal tube					
ISO 11990-1	2011-08	Ν	shaft					
			Lasers and laser-related equipment					
			Determination of laser resistance of					
			tracheal tubes Part_2: Tracheal tube					
ISO 11990-2	2010-07	Ν	cuffs					
			Health informatics Digital imaging and					
			communication in medicine (DICOM)					
ISO 12052	2006-11	N	including workflow and data management Acoustics Procedures for the					
			measurement of real-ear acoustical	Y	NO	ABNT NBR ISO	2010	
ISO 12124	2001-03	Ν	characteristics of hearing aids	I	NO	ABINT INDICISO	2010	
100 12121	2001.00		Implants for surgery Mechanical					
			testing of implantable spinal devices					
			Fatigue test method for spinal implant	Y	NO	ABNT NBR ISO	2006	
ISO 12189	2008-05	N	assemblies using an anterior support					
			Medical gloves made from natural					
			rubber latex Determination of water-					
			extractable protein using the modified	Y	NO	ABNT NBR ISO	2010	
ISO 12243	2003-10	N	Lowry method					
			Tissue paper and tissue products Part_1:	V	NO		2012	
ISO 12625-1	2011-08	Ν	General guidance on terms	Y	NO	ABNT NBR ISO	2012	

	1							
ISO 12625-12	2010-01	N	Tissue paper and tissue products Part_12: Determination of tensile strength of perforated lines Calculation of perforation efficiency	Y	NO	ABNT NBR ISO	2012	
ISO 12625-3	2005-04	N	Tissue paper and tissue products Part_3: Determination of thickness, bulking thickness and apparent bulk density	Y	NO	ABNT NBR ISO	2010	
ISO 12625-4	2005-04	N	Tissue paper and tissue products Part_4: Determination of tensile strength, stretch at break and tensile energy absorption	Y	NO	ABNT NBR ISO	2010	
			Tissue paper and tissue products Part_5:					
ISO 12625-5	2005-04	N	Determination of wet tensile strength					
100 40005 0	0005.00	N	Tissue paper and tissue products Part_6:					
ISO 12625-6	2005-02	N	Determination of grammage Tissue paper and tissue products Part_7:					
ISO 12625-7	2007-03	Ν	Determination of optical properties					
ISO 12625-8	2010-12	Ν	Tissue paper and tissue products Part_8: Water-absorption time and water-absorption capacity, basket-immersion test method	Y	NO	ABNT NBR ISO	2011	
ISO 12625-9	2005-05	N	Tissue paper and tissue products Part_9: Determination of ball burst strength	Y	NO			
ISO 12864	1997-12	N	Ophthalmic optics Contact lenses Determination of scattered light	Y	NO	ABNT NBR ISO	2010	
ISO 12865	2006-07	Ν	Ophthalmic instruments Retinoscopes					
ISO 12866	1999-06	N	Ophthalmic instruments Perimeters					
ISO 12866 AN		N	Ophthalmic instruments Perimeters; Amendment_1					
ISO 12867	2010-06	N	Ophthalmic instruments Trial frames					
ISO 12870	2004-08	N	Ophthalmic optics Spectacle frames Requirements and test methods					
ISO 12891-1	2011-05	N	Implants for surgery Retrieval and analysis of surgical implants Part_1: Retrieval and handling					
ISO 12891-2	2000-02	N	Retrieval and analysis of surgical implants Part_2: Analysis of retrieved metallic surgical implants					
130 12091-2	2000-02	N	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	N	ceramic surgical implants					
			Health informatics Service architecture					
ISO 12967-1	2009-08	N	Part_1: Enterprise viewpoint					
			Health informatics Service architecture	Y	NO	ABNT NBR ISO	2002	
ISO 12967-2	2009-08	N	Part_2: Information viewpoint	-				
100 40007 0	2009-08	Ν	Health informatics Service architecture Part_3: Computational viewpoint					
ISO 12967-3	2009-08	IN						
			Ophthalmic optics Contact lens care products Guidelines for determination					
100 40040	2011.05	N	of shelf-life					
ISO 13212 ISO 13294	2011-05 1997-05	N N	Dental handpieces Dental air-motors	Y	Y			
130 13294	1997-05	IN	Dental handpieces Dental all-motors	ř	ř			
ISO 13295	2007-07	N	Dentistry Mandrels for rotary instruments					
			Implants for surgery Ceramic					
			materials based on yttria-stabilized					
ISO 13356	2008-06	N	tetragonal 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:					
ISO 13397-2	2005-06	N	Periodontal curettes of Gr-type					
			Periodontal curettes, dental scalers and excavators Part_3: Dental scalers H-					
ISO 13397-3	1996-09	Ν	type					
	1000 00		Periodontal curettes, dental scalers and					
			excavators Part_4: Dental excavators					
ISO 13397-4	1997-12	Ν	Discoid type					
			Surgical and dental hand instruments					
00 10 100	1005.00	<u>.</u>	Determination of resistance against					
ISO 13402	1995-08	N	autoclaving, corrosion and thermal exposure Prosthetics and orthotics Categorization					
			and description of external orthoses and					
ISO 13404	2007-07	Ν	orthotic components					
	2007 07		Prosthetics and orthostics Classification					
			and description of prosthetic components					
			Part_1: Classification of prosthetic					
ISO 13405-1	1996-10	Ν	components					

		Dreathatics and arthastics Classification					
1006 10	N						
1996-10	IN						
1006 10	N						
1990-10	IN						
2008-06	N						
2003-03	N						
		Aseptic processing of health care					
2006-09	Ν	products - Part 3: Lyophilization					
2005-11	N						
2003-11		technologies					
0000 44							
2006-11	N						
			Y	NO	ABNT NBR ISO	2006	
2005-06	N	products Part_6: Isolator systems	-				
		Medical devices Quality management	v	NO		2006	
		systems Requirements for regulatory	I I	NO	ABINT NBIC ISO	2000	
2003-07	N	purposes					
		Medical devices - Quality management					
			Y	NO	ABNT NBR ISO	2006	
2000-08	N						
2009-00	IN						
		Health informatics - Electronic health record					
2008-02	Ν						
					<u> </u>	1	
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2008-12	Ν						
				1		1	
1							
2009-02	Ν						
1		communication Part_5: Interface					
2010-03	Ν	specification					
	2009-08 2008-02 2008-12 2009-02	1996-10 N 2008-06 N 2003-03 N 2006-09 N 2005-11 N 2006-11 N 2005-06 N 2003-07 N 2003-07 N 2009-08 N 2008-12 N 2009-02 N	Prosthetics and orthostics Classification and description of prosthetic components Part_3: Description of upper-limb prosthetic components 1996-10 N components 2008-06 N requirements 2008-06 N requirements 2003-03 N products Part_1: General requirements 2006-09 N products Part_2: Filtration 2006-09 N products Part_3: Lyophilization Aseptic processing of health care products Part_4: Clean-in-place products Part_4: Clean-in-place 2005-11 N technologies 2006-11 N products Part_5: Sterilization in place 2005-06 N products Part_6: Isolator systems 2005-06 N products Part_6: Isolator systems 2003-07 N purposes 2003-07 N purposes 2008-02 N Medical devices Quality management systems Requirements for regulatory purposes; Technical Corrigendum_1 2008-02 N murposes; Technical Corrigendum_1 2008-02 N murposes; Technical Corrigendum_1 2008-12 N	and description of prosthetic components Part_2: Description of lower-limb prosthetic components 1996-10 N Prosthetics and orthostics Classification and description of prosthetic components Part_3: Description of upper-limb prosthetic components 1996-10 N Aseptic processing of health care products Part_1: General 2008-06 N requirements 2003-03 N products Part_2: Filtration Aseptic processing of health care products Part_3: Lyophilization Aseptic processing of health care products Part_4: Clean-in-place 2005-09 N products Part_5: Sterilization in place 2005-11 N Aseptic processing of health care products Part_6: Isolator systems 2005-06 N Products Part_6: Isolator systems 2005-06 N Products Part_6: Isolator systems Y 2003-07 N Purposes Medical devices Quality management systems Requirements for regulatory purposes; Technical Corrigendum_1 2008-02 N Purposes; Technical Corrigendum_1 Y 2008-02 N	and description of prosthetic components Part_2: Description of lower-limb prosthetic 1996-10 N components and description of prosthetic components Part_3: Description of upper-limb prosthetic 1996-10 N components prosthetic sand orthostics Classification and description of upper-limb prosthetic 1996-10 N components products Part_1: General 2008-06 N requirements 2003-03 N products Part_2: Filtration 2006-09 N products Part_3: Lyophilization 2005-01 N technologies 2005-11 N technologies 2005-06 N products Part_4: Clean-in-place 2005-06 N products Part_5: Sterilization in place 2005-06 N products Part_6: Isolator systems 2005-06 N products Part_6: Isolator systems 2003-07 N purposes Medical devices Quality management systems Requirements for regulatory 2009-08 N purposes; Technical Corrigendum_1 2008-02 N communication Part_1: Reference 2009-08 N<	and description of prosthetic components Part 2: Description of lower-limb prosthetic 1996-10 N 2008-06 N requirements 2008-06 N 2008-06 N requirements 2008-07 N products_ Part_2: Elitration 2006-09 N products_ Part_4: Clean-in-place 2006-11 N technologies 2005-11 N technologies 2005-06 N products Part_5: Sterilization in place 2005-06 N products Part 6: Isolator systems 2005-06 N products Part 1: Requirements for regulatory ystems Requirements for regulatory 2009-08 N purposes <tr< td=""><td>and description of prosthetic components Part, 2: Description of lower-limb prosthetic components - 1998-10 N components 2008-06 N requirements 2008-06 N requirements 2008-06 N requirements 2008-07 N products Part, 2: Filtration 2006-09 N Aseptic processing of health care products Part, 3: Lyophilization - 2006-11 N products Part, 3: Lyophilization 2006-11 N products Part, 3: Clean-in-place 2006-11 N products Part, 5: Sterilization in place 2005-06 N products Part, 6: Isolator systems 2005-06 N products Part, 6: Isolator systems 2003-07 N purposes 2009-08 N purposes; Technical Corrigendum_1 2008-02 N Medical devices Quality management systems Requirements for regulatory purposes. 2008-02 N purposes; Technical Corrigendum_1 2</td></tr<>	and description of prosthetic components Part, 2: Description of lower-limb prosthetic components - 1998-10 N components 2008-06 N requirements 2008-06 N requirements 2008-06 N requirements 2008-07 N products Part, 2: Filtration 2006-09 N Aseptic processing of health care products Part, 3: Lyophilization - 2006-11 N products Part, 3: Lyophilization 2006-11 N products Part, 3: Clean-in-place 2006-11 N products Part, 5: Sterilization in place 2005-06 N products Part, 6: Isolator systems 2005-06 N products Part, 6: Isolator systems 2003-07 N purposes 2009-08 N purposes; Technical Corrigendum_1 2008-02 N Medical devices Quality management systems Requirements for regulatory purposes. 2008-02 N purposes; Technical Corrigendum_1 2

			Ophthalmic optics Spectacle lenses					
ISO 13666	1998-08	N	Vocabulary					
			Dentistry Reversible-irreversible					
ISO 13716	1999-05	N	hydrocolloid impression material systems					
ISO 13779-1	2008-10	N	Implants for surgery Hydroxyapatite Part_1: Ceramic hydroxyapatite					
ISO 13779-2	2008-10	N	Implants for surgery Hydroxyapatite Part_2: Coatings of hydroxyapatite					
ISO 13779-3	2008-02	N	Implants for surgery Hydroxyapatite Part_3: Chemical analysis and characterization of crystallinity and phase purity					
130 13779-3	2006-02	IN						
ISO 13779-4	2002-05	N	Implants for surgery Hydroxyapatite Part_4: Determination of coating adhesion strength					
130 13779-4	2002-05	IN	Poly(L-lactide) resins and fabricated					
ISO 13781	1997-02	N	forms for surgical implants In vitro degradation testing	Y	NO	ABNT NBR ISO	2000	
			Implants for surgery Metallic materials Unalloyed tantalum for					
ISO 13782	1996-12	N	surgical implant applications					
ISO 13897	2003-02	N	Dentistry Amalgam capsules					
ISO 13897 Tech	nr 2003-12	N	Dentistry Amalgam capsules; Technical Corrigendum_1					
ISO 13926-1	2004-11	N	Pen systems Part_1: Glass cylinders for pen-injectors for medical use					
ISO 13926-2	2011-04	N	Pen systems Part_2: Plunger stoppers for pen-injectors for medical use					
ISO 13958	2009-04	N	Concentrates for haemodialysis and related therapies					
ISO 13959	2009-04	N	Water for haemodialysis and related therapies					
ISO 13960	2010-07	N	Cardiovascular implants and extracorporeal systems Plasmafilters					

			Oliniaal investigation of medical devices				T	
			Clinical investigation of medical devices		5			DD0 00/0000
			for human subjects Good clinical		Р			RDC 69/2009
ISO 14155	2011-02	N	practice					
			Clinical investigation of medical devices					
			for human subjects Good clinical					
ISO 14155 Teo	c 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					
			control of a sterilization process for					
ISO 14160	2011-07	N	medical devices					
			Sterilization of health care products					
			Biological indicators Guidance for the					
			selection, use and interpretation of					
ISO 14161	2009-09	N	results					
ISO 14233	2003-03	N	Dentistry Polymer-based die materials					
			Implants for surgery Wear of total hip-					
			joint prostheses Part_1: Loading and	Y	NO	ABNT NBR ISO	2003	
			displacement parameters for wear-	I	NO	ABIT NBR 150	2003	
			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					

			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					
100 4 40 40 0	0000 11	N	knee-joint prostheses Part_2:					
ISO 14243-2	2009-11	N	Methods of measurement					
			Implants for surgery Wear of total					
			knee-joint prostheses Part_3: Loading					
			and displacement parameters for wear-					
			testing machines with displacement					
			control and corresponding					
ISO 14243-3	2004-09	N	environmental conditions 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					
ISO 14243-3		N	environmental conditions for test					
ISO 14356	2003-03	N	Dentistry Duplicating material					
			Tracheal tubes designed for laser surgery Requirements for marking					
ISO 14408	2005-06	N	and accompanying information					
100 14400	2003-00		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	N	requirements					
			Non-active surgical implants					
			Mammary implants Particular	Y	Y			
ISO 14607	2007-02	N	requirements			ABNT NBR ISO	2013	RDC 16/2012
180 14620	2008.01	N	Non-active surgical implants General					
ISO 14630	2008-01	N	requirements					

			Implants for surgery Active					
			implantable medical devices Part_1:					
			General requirements for safety,					
			marking and for information to be					
ISO 14708-1	2000-11	N	provided by the manufacturer					
	2000 11		Implants for surgery Active					
			implantable medical devices Part_2:					
ISO 14708-2	2005-10	N	Cardiac pacemakers					
100 147 00-2	2003-10		Implants for surgery Active					
			implantable medical devices Part_3:					
ISO 14708-3	2008-11	N	Implantable neurostimulators					
130 14706-3	2000-11	IN	Implants for surgery Active					
			implantable medical devices Part_4:					
ISO 14708-4	2008-11	N	Implantable infusion pumps					
130 14708-4	2000-11		Implants for surgery Active					
			implantable medical devices Part_5:	Y	NO	ABNT NBR ISO	2003	
ISO 14708-5	2010-02	N	Circulatory support devices	I	NO	ABINT INBR 130	2003	
150 14706-5	2010-02	IN						
			Implants for surgery Active					
			implantable medical devices Part_6:					
			Particular requirements for active					
			implantable medical devices intended to treat tachyarrhythmia (including					
ISO 14708-6	2010-03	N	implantable defibrillators)					
150 14708-0	2010-03	N	Ophthalmic optics Contact lens care					
			products Microbiological requirements	Y	NO	ABNT NBR ISO	2010	
			and test methods for products and	ř	NO	ABINT INDR 150	2010	
100 1 1700	2001-04	N	regimens for hygienic management of					
ISO 14729	2001-04	N	contact lenses					
			On hith almia antica. Constant lana ann					
			Ophthalmic optics Contact lens care products Microbiological requirements					
			and test methods for products and					
	10010 10	N	regimens for hygienic management of					
ISO 14729 AN	12010-10	N	contact lenses; Amendment_1					
			Ophthalmic optics Contact lens care					
			products Antimicrobial preservative					
100 4 4700	0000 00		efficacy testing and guidance on					
ISO 14730	2000-09	N	determining discard date Dentistry Implants Dynamic fatigue test					
ISO 14801	2007-11	Ν	for endosseous dental implants					
130 14001	2007-11	IN						

ISO 14879-1	2000-06	N	Implants for surgery Total knee-joint prostheses Part_1: Determination of endurance properties of knee tibial trays	Y	NO	ABNT NBR ISO	2009	
			Ophthalmic optics Spectacle lenses					
			Fundamental requirements for uncut					
ISO 14889	2003-05	Ν	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					
ISO 14937	2009-10	N	for medical devices					
			Implants for surgery Two-part addition-	Y	Y			
ISO 14949	2001-10	Ν	cure silicone elastomers	T	T	ABNT NBR ISO	2011	RDC 16/2012
			Medical devices Application of risk	Y	NO	ABNT NBR ISO	2008	
ISO 14971	2007-03	N	management to medical devices	I	NO		2000	
			Sterile obturators for single use with					
			over-needle peripheral intravascular	Y	NO	ABNT NBR ISO	2008	
ISO 14972	1998-12	N	catheters					
			Anaesthetic and respiratory equipment					
ISO 15001	2010-06	N	Compatibility with oxygen					
			Flow-metering devices for connection to					
ISO 15002	2008-07	N	terminal units of medical gas pipeline					
150 15002	2008-07	IN	systems					
			Ophthalmic instruments Fundamental					
			requirements and test methods					
			Part_1: General requirements					
ISO 15004-1	2006-06	N	applicable to all ophthalmic instruments					
	2000 00							
			Ophthalmic instruments Fundamental					
			requirements and test methods					
ISO 15004-2	2007-02	N	Part_2: 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	N	units					

[Dental elevators Part_1: General				
100 15097 1	1999-11	N	requirements				
ISO 15087-1	1999-11	N	Dental elevators Part_2: Warwick James				
ISO 15087-2	2000-04	N	elevators				
ISO 15087-2	2000-04	N	Dental elevators Part_3: Cryer elevators				
100 10007-0	2000-03		Dental elevators_ Part_4: Coupland				
ISO 15087-4	2000-05	N	elevators				
ISO 15087-5	2000-05	N	Dental elevators Part_5: Bein elevators				
ISO 15087-6	2000-05	N	Dental elevators Part_6: Flohr elevators				
			Dental tweezers Part_1: General				
ISO 15098-1	1999-10	N	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				
			Self-adhesive hanging devices for				
			infusion bottles and injection vials				
ISO 15137	2005-07	N	Requirements and test methods				
			Implants for surgery Metal				
			intramedullary nailing systems Part_1:				
ISO 15142-1	2003-08	N	Intramedullary nails				
100 101 12 1	2000 00		Implants for surgery Metal				
			intramedullary nailing systems Part_2:				
ISO 15142-2	2003-08	Ν	Locking components				
100 10142-2	2003-00	IN	Implants for surgery Metal				
			intramedullary nailing systems Part_3:				
100 454 40 0	0000.00		Connection devices and reamer				
ISO 15142-3	2003-08	N	diameter measurements				
			In vitro diagnostic medical devices				
			Measurement of quantities in samples				
			of biological origin Requirements for				
			content and presentation of reference				
ISO 15193	2009-05	N	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	N	content of supporting documentation				
			In vitro diagnostic test systems				
			Requirements for blood-glucose monitoring		\ <i>`</i>		
			systems for self-testing in managing	Y	Y	There is no national	IN nº 09/13
ISO 15197	2003-05	N	diabetes mellitus			reference	

			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					
100 10220 1	2007 01							
			Medical devices Symbols to be used					
			with medical device labels, labelling and					
			information to be supplied Part_1:					
ISO 15223-1 A	2008-06	N	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	N	validation					
			Medical devices Quality					
			management Medical device					
ISO 15225	2010-05	N	nomenclature data 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					
ISO 15254	2009-07	N	low vision					
			Implants for surgery Requirements for					
ISO 15374	1998-08	N	production of forgings					
			Medical infusion bottles Suspension					
			devices for multiple use					
ISO 15375	2010-06	N	Requirements and test methods					
			Primary packaging materials for medicinal					
			products Particular requirements for the application of ISO_9001:2008, with	Y	NO	ABNT NBR ISO	2009	
			reference to Good Manufacturing	T	NO	ABINT INDR ISO	2009	
ISO 15378	2011-11	Ν	Practice_(GMP)					
	-		Dental handpieces Air-powered scalers					
ISO 15606	1999-12	N	and scaler tips					
			Urine-absorbing aids General guidelines					
ISO 15621	2011-02	N	on evaluation					
ISO 1563	1990-09	N	Dental alginate impression material					

			Dental aqueous impression materials based					
ISO 1564	1995-11	N	on agar					
			Cardiovascular implants and artificial					
			organs Hard-shell cardiotomy/venous					
			reservoir systems (with/without filter)					
ISO 15674	2009-04	N	and soft venous reservoir bags					
	2000 01		Cardiovascular implants and artificial					
			organs Cardiopulmonary bypass					
ISO 15675	2009-04	Ν	systems Arterial blood line filters					
	2000 01		Cardiovascular implants and artificial					
			organs Requirements for single-use					
			tubing packs for cardiopulmonary					
			bypass and extracorporeal membrane					
ISO 15676	2005-07	N	oxygenation (ECMO)					
130 13070	2003-07	IN	Plastic containers for intravenous					
ISO 15747	2010-04	N	injections					
130 13747	2010-04	IN	Ophthalmic instruments					
			Endoilluminators					
			requirements and test methods for					
ISO 15752	2010-01	Ν	optical radiation safety					
150 157 52	2010-01	11	Medical infusion equipment Plastics					
			caps with inserted elastomeric liner for					
			containers manufactured by the blow-fill-					
ISO 15759	2005-04	Ν	seal (BFS) process					
150 157 59	2003-04		Ophthalmic implants Ophthalmic			Revision Project ABNT NBR		
ISO 15798	2010-01	Ν	viscosurgical devices	Y	NO	ISO	2013	
130 137 90	2010-01	11	Implants for surgery Copolymers and			100		
			blends based on polylactide In vitro	Y	NO	ABNT NBR ISO	2011	
ISO 15814	1999-11	Ν	degradation testing	I	NO	ABINT INDICISO	2011	
ISO 15814	2006-10	N	Dentistry Wires for use in orthodontics					
ISO 15854	2005-07	N	Dentistry Casting and baseplate waxes					
			Sterilization of health care products					
			Chemical indicators Guidance for					
			selection, use and interpretation of					
ISO 15882	2008-09	Ν	results					
			Washer-disinfectors Part_1: General					
			requirements, terms and definitions and					
ISO 15883-1	2006-04	Ν	tests					
	_000 01							

	1							
ISO 15883-2	2006-04	N	Washer-disinfectors Part_2: Requirements and tests for washer- disinfectors employing thermal disinfection for surgical instruments, anaesthetic equipment, bowls, dishes, receivers, utensils, glassware, etc.	Y	Y	ABNT NBR ISO 15883-2	2013	IN nº 09/13
ISO 15883-3	2006-04	N	Washer-disinfectors Part_3: Requirements and tests for washer- disinfectors employing thermal disinfection for human waste containers					
ISO 15883-4	2008-05	N	Washer-disinfectors Part_4: Requirements and tests for washer- disinfectors employing chemical disinfection for thermolabile endoscopes	Y	Y	There is no national reference		IN nº 09/13
			Washer-disinfectors Part_6: Requirements and tests for washer- disinfectors employing thermal disinfection for non-invasive, non-critical medical devices and healthcare	Y	Y	There is no national		IN nº 09/13
ISO 15883-6	2011-04	N	equipment			reference		
ISO 15912	2006-10	Ν	Dentistry Casting investments and refractory die materials					
			Dentistry Casting investments and refractory die materials; Amendment_1: Requirement and test method for adequacy of expansion of Type_1 and Type_2					
ISO 15912 AMD	2011-07	N	materials Urine-absorbing aids Basic principles for evaluation of single-use adult-incontinence- absorbing aids from the perspective of users					
ISO 16021	2000-11	N	and caregivers					
ISO 16034	2002-02	N	Ophthalmic optics Specifications for single-vision ready-to-wear near-vision spectacles					
ISO 16034 Tec	2006-08	N	Ophthalmic optics Specifications for single-vision ready-to-wear near- vision spectacles; Technical Corrigendum_1					

-			Rubber condoms for clinical trials					
ISO 16037	2002-05	N	Measurement of physical properties					
130 10037	2002-03	IN	Rubber condoms for clinical trials					
			Measurement of physical properties;					
ISO 16037 AMD	2011-02	N	Amendment 1					
100 10007 / 100	2011 02		Rubber condoms Guidance on the use of					
			ISO_4074 in the quality management of					
ISO 16038	2005-11	Ν	natural rubber latex condoms					
			Implants for surgery Minimum data					
ISO 16054	2000-12	N	sets for surgical implants					
			Dentistry Required elements for					
ISO 16059	2007-08	N	codification 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	Y	NO	ABNT NBR ISO	2008	
ISO 16201	2006-10	N	systems for daily living	-				
100 10201	2000 10		Ophthalmic optics Information					
			interchange for ophthalmic optical					
ISO 16284	2006-03	N	equipment					
100 10204	2000-03	11	Aids for ostomy and incontinence					
			Irrigation sets Requirements and test					
ISO 16391	2002-10	Ν	methods					
			Implants for surgery Acrylic resin					
			cement Flexural fatigue testing of					
			acrylic resin cements used in					
ISO 16402	2008-05	Ν	orthopaedics					
	2000 00		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	2010-02	N	interdental brushes; Amendment_1					
			Implants for surgery Test solutions					
			and environmental conditions for static					
			and dynamic corrosion tests on					
			implantable materials and medical					
ISO 16428	2005-04	N	devices					

			Implants for surgery Measurements of					
			open-circuit potential to assess					
			corrosion behaviour of metallic					
			implantable materials and medical					
100 10100	2004 07	N						
ISO 16429	2004-07	N	devices over extended time periods					
			Tracheobronchial tubes Sizing and					
ISO 16628	2008-11	N	marking					
			Ophthalmic implants Irrigating					
ISO 16671	2003-05	N	solutions for ophthalmic surgery					
			Ophthalmic implants Ocular					
ISO 16672	2003-02	Ν	endotamponades					
			Wheelchair seating Part_1: Vocabulary,					
			reference axis convention and measures for					
			body segments, posture and postural					
ISO 16840-1	2006-03	Ν	support surfaces					
			Wheelchair seating Part_2: Determination					
			of physical and mechanical characteristics					
			of devices intended to manage tissue					
ISO 16840-2	2007-07	N	integrity Seat cushions					
			Wheelchair seating Part_3: Determination					
			of static, impact and repetitive load					
ISO 16840-3	2006-07	N	strengths for postural support devices					
			Wheelchair seating Part_4: Seating	Y	NO	ABNT NBR ISO	2008	
ISO 16840-4	2009-03	N	systems for use in motor vehicles	•	110		2000	
			Health informatics Public key					
			infrastructure Part_1: Overview of digital	Y	NO	ABNT NBR ISO	2008	
ISO 17090-1	2008-02	N	certificate services					
100 17000 0			Health informatics Public key	Y	NO	ABNT NBR ISO	2008	
ISO 17090-2	2008-02	N	infrastructure Part_2: Certificate profile		_			
			Health informatics Public key					
100 47000 0	0000.00		infrastructure Part_3: Policy management					
ISO 17090-3	2008-02	N	of certification authority Health informatics Vocabulary for					
ISO 17115	2007-07	N	terminological systems					
150 17 115	2007-07	N	Urine-absorbing aids for incontinence Test					
			methods for characterizing polymer-based					
			absorbent materials Part_1: Determination					
ISO 17190-1	2001-12	N	of pH					
150 17 190-1	2001-12	IN	Urine-absorbing aids for incontinence Test					
			methods for characterizing polymer-based					
			absorbent materials Part_10:					
			Determination of extractable polymer					
ISO 17190-10	2001-12	N	content by potentiometric titration					
100 11 100-10	2001-12	IN	ounterit by potentionethe infation		1		I	

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			Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_11:			
			Determination of content of respirable			
ISO 17190-11	2001-12	N	particles			
			Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_2: Determination			
ISO 17190-2	2001-12	N	of amount of residual monomers			
130 17 190-2	2001-12	IN	Urine absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_3: Determination			
			of particle size distribution by sieve			
ISO 17190-3	2001-12	N	fractionation			
130 17 190-3	2001-12	IN	Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_4: Determination			
			of moisture content by mass loss upon			
ISO 17190-4	2001-12	Ν	heating			
130 17 130-4	2001-12	IN	Urine-absorbing aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_5: Gravimetric			
			determination of free swell capacity in saline			
ISO 17190-5	2001-12	N	solution			
	2001 12					
			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	Ν	saline solution after centrifugation			
			Ŭ l			
			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			
			Urine-absorping aids for incontinence Test			
			methods for characterizing polymer-based			
			absorbent materials Part_8: Gravimetric			
ISO 17190-8	2001-12	N	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			

r			Urine-absorbing aids for incontinence Test		1	
			methods for characterizing polymer-based			
			absorbent materials Part_9: Gravimetric			
			determination of density; Technical			
ISO 17190-9 Te	ec 2002-10	N	Corrigendum_1			
			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			
150 17 191	2004-02	11	Health informatics Messages and			
			communication Web access to DICOM			
ISO 17432	2004-12	Ν	persistent objects			
			Sleep apnoea breathing therapy			
			Part_1: Sleep apnoea breathing therapy			
ISO 17510-1	2007-10	N	equipment			
			Sleep apnoea breathing therapy			
			Part_2: Masks and application			
ISO 17510-2	2007-10	N	accessories			
			In vitro diagnostic medical devices			
			Measurement of quantities in biological			
			samples Metrological traceability of			
			values assigned to calibrators and			
ISO 17511	2003-08	N	control materials			
			Clinical laboratory testing and in vitro			
			medical devices Requirements for in			
			vitro monitoring systems for self-testing			
ISO 17593	2007-04	N	of oral anticoagulant therapy			
			Sterilization of medical devices			
			Information to be provided by the			
			manufacturer for the processing of			
ISO 17664	2004-03	N	resterilizable medical devices			
			Sterilization of health care products			
			Moist heat Part_1: Requirements for			
			the development, validation and routine			
			control of a sterilization process for			
ISO 17665-1	2006-08	N	medical devices			
			Wear of implant materials Polymer			
100 47050			and metal wear particles Isolation and			
ISO 17853	2011-03	N	characterization			
ISO 1797-1	2011-08	N	Dentistry Shanks for rotary instruments Part_1: Shanks made of metals			
130 1797-1	2011-00	IN	rait_1. Shahks made of metals			

			Dental rotary instruments; shanks; part_2:			
ISO 1797-2	1992-02	Ν	shanks made of plastics			
ISO 18084	2011-09	N	Press tools for tablets Punches and dies			
			Health informatics Integration of a			
ISO 18104	2003-12	N	reference terminology model for nursing			
			In vitro diagnostic medical devices			
			Information supplied by the			
			manufacturer (labelling) Part_1:			
			Terms, definitions and general			
ISO 18113-1	2009-12	N	requirements			
			In vitro diagnostic medical devices			
			Information supplied by the			
			manufacturer (labelling) Part_2: In			
			vitro diagnostic reagents for			
ISO 18113-2	2009-12	N	professional use			
			In vitro diagnostic medical devices			
			Information supplied by the			
			manufacturer (labelling) Part_3: In			
			vitro diagnostic instruments for			
ISO 18113-3	2009-12	N	professional use			
			In vitro diagnostic medical devices			
			Information supplied by the			
			manufacturer (labelling) Part_4: In			
ISO 18113-4	2009-12	N	vitro diagnostic reagents for self-testing			
			In vitro diagnostic medical devices			
			Information supplied by the			
			manufacturer (labelling) Part_5: In			
			vitro diagnostic instruments for self-			
ISO 18113-5	2009-12	N	testing			
			In vitro diagnostic medical devices			
			Measurement of quantities in biological			
			samples Metrological traceability of			
			values for catalytic concentration of			
			enzymes assigned calibrators and			
ISO 18153	2003-08	N	control materials			

			Implants for surgery Wear of total					
			intervertebral spinal disc prostheses					
			Part_1: Loading and displacement					
			parameters for wear testing and					
			corresponding environmental conditions					
ISO 18192-1	2011-03	N	for test					
100 10102 1	2011 00		Implants for surgery Wear of total					
			intervertebral spinal disc prostheses					
ISO 18192-2	2010-06	Ν	Part_2: Nucleus replacements					
100 10102 2	2010 00		Health Informatics Messages and					
			communication Format of length limited	Y	NO	ABNT NBR ISO	2010	
ISO 18232	2006-04	N	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 and recommendations for					
ISO 18369-1	2006-08	N	labelling specifications					
			Ophthalmic optics Contact lenses					
			Part_1: Vocabulary, classification					
			system and recommendations for					
ISO 18369-1 A	12009-02	N	labelling specifications; Amendment_1					
			Ophthalmic optics Contact lenses					
ISO 18369-2	2006-08	N	Part_2: Tolerances					
			Ophthalmic optics Contact lenses					
ISO 18369-3	2006-08	N	Part_3: Measurement methods					
			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					
ISO 18472	2006-06	N	Test equipment					
			The second ship for the second second second					
100 40777	0005 00		Transportable liquid oxygen systems for					
ISO 18777	2005-02	N	medical use Particular requirements					
100 40770	0005 00	N 1	Respiratory equipment Infant					
ISO 18778	2005-02	N	monitors Particular requirements					
			Medical devices for conserving oxygen					
			and oxygen mixtures Particular					
ISO 18779	2005-02	N	requirements					

			Health informatics Clinical analyser	
1			interfaces to laboratory information	
ISO 18812	2003-03	Ν	systems Use profiles	
			In vitro diagnostic medical devices	
			Information supplied by the	
			manufacturer with in vitro diagnostic	
ISO 19001	2002-11	N	reagents for staining in biology	
			Rail systems for supporting medical	
ISO 19054	2005-07	Ν	equipment	
ISO 1942	2009-12	N	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	
			Dentistry Manual toothbrushes General	
ISO 20126	2012-01	Ν	requirements and test methods	
			Dentistry Powered toothbrushes	
ISO 20127	2005-03	N	General requirements and test methods	
			Implants for surgery Metallic	
			materials Classification of	
			microstructures for alpha+beta titanium	
ISO 20160	2006-05	N	alloy bars	
			Health informatics Health cards General	
ISO 20301	2006-11	N	characteristics	
			Health informatics Health cards	
ISO 20302	2006-12	N	Numbering system and registration procedure for issuer identifiers	
150 20302	2006-12	N		
			Clinical loberatory testing and in vitre	
			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	
ISO 20776-1	2006-11	N	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:			
ISO 20795-1	2008-08	N	Denture base polymers			
			Dentistry Base polymers Part_1:			
ISO 20795-1 Te	~ 2000 02	N	Denture base polymers; Technical Corrigendum_1			
130 20795-1 16	ed 2009-02	IN	Dentistry Base polymers Part_2:			
ISO 20795-2	2010-03	Ν	Orthodontic base polymers			
			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			
100 20001	2010 00		Health informatics Harmonized data types			
ISO 21090	2011-02	Ν	for information interchange			
			Medical gloves Determination of			
ISO 21171	2006-05	Ν	removable surface powder			
			Dentistry Materials used for dental			
			equipment surfaces Determination of			
ISO 21530	2004-06	N	resistance to chemical disinfectants			
			Dentistry Graphical symbols for dental			
ISO 21531	2009-02	N	instruments			
100 04 500	0000.00	N	Dentistry Reusable cartridge syringes			
ISO 21533	2003-06	N	intended for intraligamentary injections Dentistry Reusable cartridge syringes			
			intended for intraligamentary injections;			
ISO 21533 Tec	br 2009-12	Ν	Technical Corrigendum_1			
2021000100			Non-active surgical implants Joint			
			replacement implants Particular			
ISO 21534	2007-10	N	requirements			
100 21004	2007 10		Non-active surgical implants Joint			
			replacement implants Specific			
			requirements for hip-joint replacement			
ISO 21535	2007-10	Ν	implants			
100 21000	2007-10	IN	Impianto			L

1		Non active surgical implants loint					T
2007-10	N						
2004-05	N						
0004.05	N						
2004-05	N						
2004 05	N						
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2008-04	N						
2000-04	IN						
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2000-04							
2007-06	N						
2001 00							
2010-06	Ν						
1992-06	Ν						
2007-06	N	orthodontics					
		Needle-free injectors for medical use -					
2006-06	N	-					
2000 00							
2010-12	Ν						
2011-04	N	Dentistry Rotary polishers; Amendment_1					
		Dentistry Periodontal probes Part_1:					
2012-04	N	General requirements					
		High-pressure flexible connections for					
2009-10	N	0 1					
2009-10	N						
2000 10							
2005-11	N						
2000 11							
2005-08	N		Y	NO			
					1		
2005-09	Ν	powered scalers and scaler tips	Y	NO	1	1	1
	2007-06 2006-06 2010-12 2006-07 2011-04 2012-04 2009-10 2009-10 2005-11 2005-08	2004-05 N 2004-05 N 2004-05 N 2006-11 N 2008-04 N 2008-04 N 2008-04 N 2007-06 N 2007-06 N 2007-06 N 2007-06 N 2007-06 N 2007-06 N 2006-06 N 2006-07 N 2011-04 N 2012-04 N 2009-10 N 2005-11 N 2005-08 N	N Health informatics Patient healthcard 2004-05 N data Part_1: General structure 2004-05 N data Part_2: Common objects 2004-05 N data Part_2: Common objects 2004-05 N data Part_3: Limited clinical data 2006-01 N data Part_5: Identification data 2008-04 N data Part_6: Administrative data 2008-04 N data Part_7: Medication data 2007-06 N data Part_8: Links 2007-06 N data Part_8: Links 2007-06 N data Part_8: Links 2007-06 N detaeth informatics Health indicators 2007-06 N orthodontics 2006-06 N Requirements and test methods 2010-12 N <td< td=""><td>replacement implants Specific requirements for knee-joint replacement 2007-10 N implants 2004-05 N dataPart_1: General structure 2004-05 N dataPart_2: Common objects Health informatics Patient healthcard 2004-05 N dataPart_2: Common objects Health informatics Patient healthcard 2006-01 N dataPart_4: Extended clinical data 2006-01 N dataPart_5: Identification data 2008-04 N dataPart_6: Administrative data 2008-04 N dataPart_6: Administrative data 2008-04 N dataPart_6: Administrative data 2008-04 Health informatics Patient healthcard 2007-06 N dataPart_8: Links Dental rotary instruments; 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			Transfer sets for pharmaceutical			1	
			preparations Requirements and test	Y	NO		
ISO 22413	2010-06	Ν	methods				
			Medical devices utilizing animal tissues				
			and their derivatives Part_1:				
ISO 22442-1	2007-12	Ν	Application of risk management				
			Medical devices utilizing animal tissues				
			and their derivatives Part_2: Controls				
ISO 22442-2	2007-12	Ν	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 encephalopathy (TSE)				
ISO 22442-3	2007-12	Ν	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 blood (fixed volume, horizontally				
ISO 22609	2004-12	Ν	projected)				
100 22000	2004 12						
			Surgical drapes, gowns and clean air				
			suits, used as medical devices, for				
			patients, clinical staff and equipment				
			Test method to determine the				
ISO 22610	2006-07	N	resistance to wet bacterial penetration				
			Clothing for protection against infectious				
			agents Test method for resistance to				
ISO 22612	2005-03	N	dry microbial penetration				
			Dentistry Metallic materials for fixed and				
ISO 22674	2006-11	N	removable restorations and appliances				
			Prosthetics Testing of ankle-foot				
			devices and foot units Requirements				
ISO 22675	2006-10	N	and test methods				
ISO 22715	2006-04	N	Cosmetics Packaging and labelling				

			Cosmetics Good Manufacturing Practices			
			(GMP) Guidelines on Good Manufacturing			
ISO 22716	2007-11	N	Practices			
100 227 10	2007-11		Dentistry Implantable materials for bone			
			filling and augmentation in oral and			
			maxillofacial surgery Contents of a			
ISO 22794	2007-07	N	technical file			
			Dentistry Membrane materials for guided			
			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	protection to facilitate trans-border nows of personal health information			
150 22037	2004-04		Implants for surgery In vitro evaluation			
			for apatite-forming ability of implant			
ISO 23317	2007-06	N	materials			
130 23317	2007-00	IN				
			Breathing system filters for anaesthetic			
100 00000 4	0000.00	N	and respiratory use Part_1: Salt test			
ISO 23328-1	2003-08	N	method to assess filtration performance			
			Breathing system filters for anaesthetic			
	0000 40	N	and respiratory use Part_2: Non-			
ISO 23328-2	2002-10	N	filtration aspects Male condoms Requirements and test			
			methods for condoms made from synthetic			
ISO 23409	2011-02	Ν	materials			
100 20 100	2011 02		Guidance for the preparation and			
			quality management of fluids for			
ISO 23500	2011-05	N	haemodialysis and related therapies			
			Assistive products for blind and vision-			
			impaired persons Tactile walking surface			
ISO 23599	2012-03	N	indicators			
			Assistive products for persons with vision			
			impairments and persons with vision and			
			hearing impairments Acoustic and tactile			
ISO 23600	2007-11	N	signals for pedestrian traffic lights			
			In vitro diagnostic medical devices			
			Evaluation of stability of in vitro			
ISO 23640	2011-12	N	diagnostic reagents			

			Anaesthetic and respiratory equipment					
			Peak expiratory flow meters for the					
			assessment of pulmonary function in					
ISO 23747	2007-07	N	spontaneously breathing humans					
			Sharps injury protection					
			Requirements and test methods					
			Sharps protection features for single-					
			use hypodermic needles, introducers for					
			catheters and needles used for blood					
ISO 23908	2011-06	N	sampling					
			Ophthalmic optics and instruments					
ISO 24157	2008-07	N	Reporting aberrations of the human eye					
ISO 24137	2008-07	N	Skin barrier for ostomy aids Vocabulary					
100 24214	2000-11		Dentistry Mercury and alloys for dental					
ISO 24234	2004-10	Ν	amalgam					
			Dentistry Mercury and alloys for dental					
			amalgam Amendment_1: Requirements					
			for marking and manufacturer's instructions					
ISO 24234 AME	2011-08	N	concerning mercury					
			Tips for assistive products for walking					
100 04445 4			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					
100 24413 2	2011-00		Ergonomics Accessible design Auditory					
ISO 24500	2010-10	Ν	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					
ISO 24503	2011-01	N	Ergonomics Accessible designTactile dots and bars on consumer products					
150 24503	2011-01	IN						
			Sterilization of medical devices Low					
			temperature steam and formaldehyde	Y	NO	ABNT NBR ISO	2008	
			Requirements for development,	I	NO	ADIVE NOR 130	2000	
			validation and routine control of a					
ISO 25424	2009-09	N	sterilization process for medical devices					

		ſ	Cardiovacaular implanta			Γ		
			Cardiovascular implants	Y	NO	ABNT NBR ISO	2008	
100 05500 4			Endovascular devices Part_1:	Y	NO	ABNT NBR ISO	2008	
ISO 25539-1	2003-03	N	Endovascular prostheses					
			Cardiovascular implants					
			Endovascular devices Part_1:	Y	NO	ABNT NBR ISO	1997	
			Endovascular prostheses;					
ISO 25539-1 A	2005-07	N	Amendment_1: Test methods					
			Cardiovascular implants					
			Endovascular devices Part_2:	Y	NO	ABNT NBR ISO	2008	
ISO 25539-2	2008-09	N	Vascular stents					
			Cardiovascular implants					
			Endovascular devices Part_3: Vena	Y	NO	ABNT NBR ISO	2008	
ISO 25539-3	2011-12	N	cava filters					
100 05700			Health informatics Genomic Sequence	Y	NO	ABNT NBR ISO	2008	
ISO 25720	2009-08	N	Variation Markup Language (GSVML) Female condoms Requirements and test		-			
ISO 25841	2011-07	Ν	methods	Y	NO	ABNT NBR ISO	2001	
100 20041	2011-07		Water treatment equipment for					
			haemodialysis applications and related	Y	NO	ABNT NBR ISO	1997	
ISO 26722	2009-04	N	therapies					
100 20122	2000 01		Anaesthetic and respiratory equipment					
			Spirometers intended for the					
			measurement of time forced expired	Y	NO	ABNT NBR ISO	1997	
ISO 26782	2009-07	N	volumes in humans					
100 20102	2000 01		Anaesthetic and respiratory equipment					
			Spirometers intended for the					
			measurement of time forced expired	Y	NO	ABNT NBR ISO	2008	
			volumes in humans; Technical					
ISO 26782 Tec	2009-11	N	Corrigendum_1					
			Anaesthetic and respiratory equipment					
			User-applied labels for syringes					
			containing drugs used during	Y	NO	ABNT NBR ISO	1999	
			anaesthesia Colours, design and					
ISO 26825	2008-08	N	performance					
100 20020	2000 00		Dentistry Brackets and tubes for use in					
ISO 27020	2010-12	Ν	orthodontics	Y	NO	ABNT NBR ISO	1998	
			Cardiac rhythm management devices					
	1		Symbols to be used with cardiac rhythm					
			management device labels, and	Y	NO	ABNT NBR ISO	1998	
			information to be supplied General					
ISO 27185	2012-02	Ν	requirements					

ISO 27186	2010-03	N	Active implantable medical devices Four-pole connector system for implantable cardiac rhythm management devices Dimensional and test requirements	Y	NO	ABNT NBR ISO	2008	
ISO 27427	2010-03	N	Anaesthetic and respiratory equipment Nebulizing systems and components	Υ	NO	ABNT NBR ISO	2004	
ISO 27799	2008-07	N	Health informatics Information security management in health using ISO/IEC_2702	Y	NO	ABNT NBR ISO	2008	
ISO 28158	2010-07	N	Dentistry Integrated dental floss and handles	Y	NO	ABNT NBR ISO	2008	
ISO 28319	2010-05	N	Dentistry Laser welding	Y	NO	Project Revision ABNT NBR ISO	2013	
ISO 28399	2011-01	N	Dentistry Products for external tooth bleaching	Y	NO	ABNT NBR ISO	2008	
ISO 28620	2010-02	N	Medical devices Non-electrically driven portable infusion devices	Y	NO	ABNT NBR ISO	2088	
ISO 29701	2010-09	N	Nanotechnologies Endotoxin test on nanomaterial samples for in vitro systems Limulus amebocyte lysate (LAL) test	Υ	NO	ABNT NBR ISO	2008	
ISO 29781	2008-12	N	Prostheses and orthoses Factors to be included when describing physical activity of a person who has had a lower limb amputation(s) or who has a deficiency of a lower limb segment(s) present at birth	Y	NO	ABNT NBR ISO	1996	
ISO 29782	2008-12	N	Prostheses and orthoses Factors to be considered when specifying a prosthesis for a person who has had a lower limb amputation	Y	NO	ABNT NBR ISO	1996	
ISO 29783-1	2008-12	N	Prosthetics and orthotics Vocabulary Part_1: Normal gait	Y	NO	ABNT NBR ISO	1997	
ISO 29941	2010-12	N	Condoms Determination of nitrosamines migrating from natural rubber latex condoms	Y	NO	ABNT NBR ISO	1997	
ISO 29942	2011-07	N	Prophylactic dams Requirements and test methods	Y	NO	ABNT NBR ISO	1998	
ISO 3107	2011-03	N	Dentistry Zinc oxide/eugenol cements and zinc oxide/non-eugenol cements	Y	NO	ABNT NBR ISO	1996	
ISO 32	1977-05	N	Gas cylinders for medical use; Marking for identification of content	Y	NO	ABNT NBR ISO	1996	

			Dentistry Root-canal instruments Part_1:				
ISO 3630-1	2008-02	Ν	General requirements and test methods				
			Dental root-canal instruments Part_2:			1	
ISO 3630-2	2000-12	Ν	Enlargers				
			Dental root-canal instruments; part_3:				
ISO 3630-3	1994-03	Ν	condensers, pluggers and spreaders				
			Dentistry Root canal instruments Part_4:				
ISO 3630-4	2009-07	N	Auxiliary instruments				
			Dentistry Endodontic instruments	Y	NO		
ISO 3630-5	2011-10	N	Part_5: Shaping and cleaning instruments				
100 0000 4	1007.00		Dental rotary instruments Burs Part_1:	Y	NO		
ISO 3823-1	1997-08	N	Steel and carbide burs				
ISO 3823-2	2003-05	N	Dentistry Rotary bur instruments Part_2: Finishing burs				
130 3023-2	2003-03	IN	Dentistry Rotary bur instruments Part_2:				
ISO 3823-2 AM	MD 2008-07	Ν	Finishing burs; Amendment_1				
			Plastics collapsible containers for				
			human blood and blood components		Y		RDC 35/2014
ISO 3826-1	2003-11	N	Part_1: Conventional containers				
100 0020 1	2000 11		Plastics collapsible containers for				
			human blood and blood components				
			Part_2: Graphical symbols for use on				
ISO 3826-2	2008-08	Ν	labels and instruction leaflets				
130 3620-2	2000-00	IN	Plastics collapsible containers for				
			human blood and blood components				
			Part_3: Blood bag systems with				
	2006-09	N					
ISO 3826-3	2006-09	N	integrated features				
			Association Defenses are faither				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_1: Reference equivalent threshold				
			sound pressure levels for pure tones				
ISO 389-1	1998-11	N	and supra-aural earphones				
			Acoustics Reference zero for the				
			calibration of audiometric equipment				
			Part_2: Reference equivalent threshold				
			sound pressure levels for pure tones				
ISO 389-2	1994-07	N	and insert earphones				

			Acquistica Deference for the					
			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					
			Acoustics Reference zero for the					
			calibration of audiometric equipment					
			Part_3: Reference equivalent treshold					
			force levels for pure tones and bone					
ISO 389-3 Tec	1995-08	Ν	vibrators; Technical corrigendum_1					
			Acoustics Reference zero for the					
			calibration of audiometric equipment					
			Part_4: Reference levels for narrow-					
ISO 389-4	1994-10	Ν	band 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					
150 389-6	2007-07	IN	Acoustics Reference zero for the					
			_					
			calibration of audiometric equipment					
			Part_7: Reference threshold of hearing					
100 000 7	0005 44		under free-field and diffuse-field					
ISO 389-7	2005-11	N	listening conditions					
			Acoustics Reference zero for the					
			calibration of audiometric equipment					
			Part_8: Reference equivalent threshold					
	0004.05		sound pressure levels for pure tones					
ISO 389-8	2004-05	N	and circumaural earphones					
			Acoustics Reference zero for the					
			calibration of audiometric equipment					
			Part_9: Preferred test conditions for the	Y	NO	ABNT NBR ISO	2013	
			determination of reference hearing					
ISO 389-9	2009-05	N	threshold levels					
	1		Dentistry Designation system for teeth and	Y	NO	ABNT NBR ISO	2013	
ISO 3950	2009-05	N	areas of the oral cavity	•			2010	
ISO 3964	1982-12	N	Dental handpieces; Coupling dimensions				+	
100 4040	2000 10	N	Dentistry Polymer-based restorative					
ISO 4049	2009-10	N	materials					

r								
			Dentistry Information system on the					
			location of dental equipment in the working					
ISO 4073	2009-07	N	area of the oral health care provider					
			Natural latex rubber condoms					
ISO 4074	2002-02	N	Requirements and test methods					
			Natural latex rubber condoms					
100 4074 Tesha	0000 40	N	Requirements and test methods; Technical	Y	Y		0010	
ISO 4074 Techn	1 2003-10	N	Corrigendum_1 Natural latex rubber condoms			ABNT NBR ISO 4074	2013	RDC 62/2008
			Requirements and test methods; Technical					
ISO 4074 Techn	2008-04	N	Corrigendum 2					
			Anaesthetic and respiratory equipment					
ISO 4135	2001-08	N	Vocabulary					
ISO 4823	2000-12	N	Dentistry Elastomeric impression materials					
	10007.07	N	Dentistry Elastomeric impression	Y	NO	ABNT NBR ISO	2010	
ISO 4823 AMD	12007-07	N	materials; Amendment_1 Dentistry Elastomeric impression					
ISO 4823 Techn	2004-07	N	materials; Technical Corrigendum_1	Y	NO	ABNT NBR ISO	2010	
100 1020 1001	200101		Anaesthetic and respiratory equipment					
			Conical connectors Part_1: Cones					
ISO 5356-1	2004-05	N	and sockets					
			Anaesthetic and respiratory equipment					
			Conical connectors Part_2: Screw-					
ISO 5356-2	2006-09	N	threaded weight-bearing connectors					
			Anaesthetic machines for use with					
ISO 5358	1992-01	N	humans					
			Low-pressure hose assemblies for use					
ISO 5359	2008-06	Ν	with medical gases					
			Low-pressure hose assemblies for use					
ISO 5359 AME	2011-12	N	with medical gases; Amendment_1					
			Anaesthetic vaporizers Agent-specific					
ISO 5360	2012-01	N	filling systems					
			Anaesthetic and respiratory equipment					
ISO 5361	1999-09	N	Tracheal tubes and connectors					
ISO 5361-4	1999-09	N	Tracheal tubes; Part 4 : Cole type					
ISO 5361-4	2006-06	N	Anaesthetic reservoir bags					
100 0002	2000-00	IN	Anaesthetic and respiratory equipment					
ISO 5364	2008-07	N	Oropharyngeal airways					
130 3304	2000-07	IN	Oropharyngear anways					

	Т		<u> </u>			<u> </u>	r1
			Anaesthetic and respiratory equipment				ļ
			Tracheostomy tubes Part_1: Tubes				Į
ISO 5366-1	2000-12	N	and connectors for use in adults		1		ļ
	1					1	l
			Anaesthetic and respiratory equipment		1		ļ
			Tracheostomy tubes Part_3:		1		ļ
ISO 5366-3	2001-08	N	Paediatric tracheostomy tubes				l
			Anaesthetic and respiratory equipment				
			Tracheostomy tubes Part_3:		1		ļ
			Paediatric tracheostomy tubes;		1		ļ
ISO 5366-3 Te	<u>¢2003-01</u>	N	Technical Corrigendum_1				ļ
100 5005	0000.00		Breathing tubes intended for use with		1		ļ
ISO 5367	2000-06	N	anaesthetic apparatus and ventilators			+	ļ
			Implants for surgery Metallic		1		ļ
100 5000 4	2007.00	N	materials Part_1: Wrought stainless		1		ļ
ISO 5832-1	2007-06	N	steel			+	ļi
			Implants for surgery Metallic		1		ļ
			materials Part_1: Wrought stainless		1		ļ
ISO 5832-1 Te	2008-04	N	steel; Technical Corrigendum_1		1		ļ
	1		Implants for surgery Metallic		ļ	+	
			materials Part_11: Wrought titanium 6-		1		ļ
ISO 5832-11	1994-09	N	aluminium 7-niobium alloy		1		ļ
			Implants for surgery Metallic			1	
			materials Part_12: Wrought cobalt-		1		ļ
ISO 5832-12	2007-05	N	chromium-molybdenum alloy				
			Implants for surgery Metallic	 			
			materials Part_12: Wrought cobalt-		1		ļ
			chromium-molybdenum alloy; Technical		1		ļ
ISO 5832-12 T	2008-09	N	Corrigendum_1			+	ļ
			Implants for surgery Metallic		1		ļ
			materials Part_14: Wrought titanium		1		
100 5000 11	0007.40		15-molybdenum 5-zirconium 3-		1		
ISO 5832-14	2007-10	N	aluminium alloy			+	
			Implants for surgery Metallia		1		ļ
ISO 5832-2	1999-07	N	Implants for surgery Metallic materials Part_2: Unalloyed titanium		1		
130 3032-2	1999-07	IN	Implants for surgery Metallic		<u> </u>	+	ŀi
			materials Part_3: Wrought titanium 6-		1		ļ
ISO 5832-3	1996-07	N	aluminium 4-vanadium alloy		1		
100 0002-0	1330-01				l		L

			Implants for surgery Metallic					
			materials Part_4: Cobalt-chromium-					
ISO 5832-4	1996-07	N	molybdenum casting alloy					
			Implants for surgery Metallic					
			materials Part_5: Wrought cobalt-	Y	NO			
ISO 5832-5	2005-10	N	chromium-tungsten-nickel alloy					
			Implants for surgery Metallic					
			materials Part_6: Wrought cobalt-					
ISO 5832-6	1997-07	N	nickel-chromium-molybdenum alloy					
			Implants for surgery; metallic materials;					
			part_7: forgeable and cold-formed					
			cobalt-chromium-nickel-molybdenum-					
ISO 5832-7	1994-02	N	iron alloy					
			Implants for surgery Metallic					
			materials Part_8: Wrought cobalt-	Y	NO	ABNT NBR ISO	2008	
			nickel-chromium-molybdenum-tungsten-	·			2000	
SO 5832-8	1997-07	N	iron alloy					
			Implants for surgery Metallic					
			materials Part_9: Wrought high	Y	NO	ABNT NBR ISO	2004	
ISO 5832-9	2007-06	N	nitrogen stainless steel					
			Implants for surgery Acrylic resin	Y	NO	ABNT NBR ISO	2012	
ISO 5833	2002-05	N	cements		_			
			Implants for surgery Ultra-high-					
			molecular-weight polyethylene Part_1:	Y	NO	ABNT NBR ISO	2011	
ISO 5834-1	2005-06	N	Powder form					
			hands at fan som som blitter blad					
			Implants for surgery Ultra-high-	Y	NO	ABNT NBR ISO	2004	
00 5004 4 7.	0007.05	N	molecular-weight polyethylene Part_1:					
ISO 5834-1 Te	2007-05	N	Powder form; Technical Corrigendum_1					
			Implants for surgery Ultra-high-	X			2002	
	2011 02	NI NI	molecular-weight polyethylene Part_2:	Y	NO	ABNT NBR ISO	2008	
ISO 5834-2	2011-08	N	Moulded forms					
			Implants for surgery Ultra-high-	Y	NO	Project Revision ABNT NBR	2013	
100 5024 2	2005 07	N	molecular-weight polyethylene Part_3:	Ť	NO	ISO	2013	
ISO 5834-3	2005-07	N	Accelerated ageing methods					
			Implanta for aurgany Ultra high					
			Implants for surgery Ultra-high- molecular-weight polyethylene Part_4:					
100 5024 4	2005-05	N	Oxidation index measurement method					
ISO 5834-4	2000-05	IN	Oxidation index measurement method		1			

					1	1	1
			Implants for surgery Ultra-high-				
			molecular-weight polyethylene Part_5:				
ISO 5834-5	2005-06	Ν	Morphology assessment method				
			Implants for surgery; metal bone screws				
			with hexagonal drive connection,				
			spherical under-surface of head,				
ISO 5835	1991-01	Ν	asymmetrical thread; dimensions				
			Implants for surgery; metal bone plates;				
			holes corresponding to screws with				
			asymmetrical thread and spherical				
ISO 5836	1988-12	Ν	under-surface				
			Implants for surgery; Intramedullary				
			nailing systems; Part 1 : Intramedullary				
			nails with cloverleaf or V-shaped cross-				
ISO 5837-1	1985-06	N	section				
			Implants for surgery; Intramedullary				
ISO 5837-2	1980-11	N	nailing systems; Part 2 : Medullary pins				
			Implants for surgery Skeletal pins and				
			wires Part_1: Material and mechanical	Y	Y		
ISO 5838-1	1995-11	N	requirements				
			Implants for surgery; skeletal pins and				
100 5000 0	1001.01	N	wires; part_2: Steinmann skeletal pins;				
ISO 5838-2	1991-01	N	dimensions				
			Implants for surgery skeletal pipe and				
190 5838-3	1003-00	Ν					
130 3030-3	1993-09						
150 5840	2005-03	Ν					
	2000 00						
ISO 5841-2	2000-10	N					
ISO 5841-3	2000-10	Ν	pacemakers				
ISO 5838-3 ISO 5840 ISO 5841-2 ISO 5841-3	1993-09 2005-03 2000-10 2000-10	N N N N	Implants for surgery; skeletal pins and wires; part_3: Kirschner skeletal wires Cardiovascular implants Cardiac valve prostheses Implants for surgery Cardiac pacemakers Part_2: Reporting of clinical performance of populations of pulse generators or leads Implants for surgery Cardiac pacemakers Part_2: Reporting of clinical performance of populations of pulse generators or leads Implants for surgery Cardiac pacemakers Part_3: Low-profile connectors [IS-1] for implantable pacemakers				

			Implants for surgery Cardiac					
			pacemakers Part_3: Low-profile					
			connectors (IS-1) for implantable					
ISO 5841-3 Te	2003-11	N	pacemakers; Technical Corrigendum_1					
	2000 11		Conical fittings with a 6 % (Luer) taper					
			for syringes, needles and certain other					
			medical equipment; Part 1 : General	Y	Y			
ISO 594-1	1986-06	N	requirements			ABNT NBR ISO	2003	C´s 03, 04 e 05/20
			Conical fittings with 6%_(Luer) taper for				2000	0 0 00, 0 1 0 00/20
			syringes, needles and certain other					
			medical equipment Part_2: Lock	Y	Y			
ISO 594-2	1998-09	N	fittings			ABNT NBR ISO	2003	C´s 03, 04 e 05/20
			Reusable all-glass or metal-and-glass					0 0 00, 0 1 0 00,24
			syringes for medical use; Part 1 :					
ISO 595-1	1986-12	N	Dimensions					
			Reusable all-glass or metal-and-glass					
			syringes for medical use; Part 2 :					
			Design, performance requirements and					
ISO 595-2	1987-12	N	tests					
			Hypodermic needles for single use;					
ISO 6009	1992-12	N	colour coding for identification		Y			RDC 05/2011
			Hypodermic needles for single use					
			Colour coding for identification;	Y	NO	ABNT NBR ISO	1997	
ISO 6009 Tech	2008-03	N	Technical Corrigendum_1					
			Dentistry Number coding system for rotary					
			instruments Part_1: General					
ISO 6360-1	2004-04	N	characteristics					
			Dentistry Number coding system for rotary instruments Part_1: General					
ISO 6360-1 Tech	2007-09	Ν	characteristics; Technical Corrigendum_1					
150 0500-1 160	2007-09		Dentistry Number coding system for rotary					
ISO 6360-2	2004-11	N	instruments Part_2: Shapes					
			Dentistry Number coding system for rotary					
			instruments Part_2: Shapes;					
ISO 6360-2 AMD	2011-12	N	Amendment_1					
			Dentistry Number coding system for rotary					
	2005 11	N	instruments Part_3: Specific					
ISO 6360-3	2005-11	N	characteristics of burs and cutters Dentistry Number coding system for rotary					
			instruments Part_4: Specific					
ISO 6360-4	2004-06	Ν	characteristics of diamond instruments					
	2004 00				Į	ļ	1	

			Dentistry Number coding system for rotary					
			instruments Part_5: Specific					
ISO 6360-5	2007-12	Ν	characteristics of root-canal instruments					
130 0300-5	2007-12	IN	Dentistry Number coding system for rotary					
			instruments Part_6: Specific					
ISO 6360-6	2004-06	Ν	characteristics of abrasive instruments					
	2004 00		Dentistry Number coding system for rotary					
			instruments Part_7: Specific					
			characteristics of mandrels and special	Y	Y			
ISO 6360-7	2006-02	Ν	instruments					
			Implants for surgery Ceramic					
			materials Part_1: Ceramic materials	Y	Y			
ISO 6474-1	2010-02	Ν	based on high purity alumina					
100 0474-1	2010-02		Implants for surgery; metal bone screws					
			with asymmetrical thread and spherical					
			under-surface; mechanical	Y	Y			
100 0475	1000 11	N	-					
ISO 6475	1989-11	N	requirements and test methods					
ISO 6710	1995-08	N	Single-use containers for venous blood specimen collection					
ISO 6872	2008-09	N N	Dentistry Ceramic materials	Y	Y			
ISO 6873	1998-03	N	Dental gypsum products	Ĭ	T			
130 0073	1990-03	IN .	Dentistry Polymer-based pit and fissure					
ISO 6874	2005-08	Ν	sealants					
100 001 4	2000 00					There is no national		
ISO 6875	2011-07	N	Dentistry Patient chair	Y	Y	reference		IN nº 09/13
ISO 6876	2001-08	N	Dental root canal sealing materials					
ISO 6877	2006-04	N	Dentistry Root-canal obturating points					
			Surgical instruments; non-cutting,					
			articulated instruments; general					
ISO 7151	1988-12	Ν	requirements and test methods					
			Surgical instruments; metallic materials;					
ISO 7153-1	1991-04	Ν	part_1: stainless steel					
	1001 01		Surgical instruments Metallic					
			materials Part_1: Stainless steel;					
ISO 7153-1 AN	1000.02	N	Amendment_1					
130 7 133-1 AN	1999-03	IN	Wheelchairs Part_1: Determination of					
ISO 7176-1	1999-10	Ν	static stability	Y	Y	ABNT NBR ISO 7176-1	2009	IN nº 09/13
	1000-10		Wheelchairs Part_10: Determination of				2005	
			obstacle-climbing ability of electrically	Y	Y	There is no national		IN nº 09/13
ISO 7176-10	2008-11	N	powered wheelchairs	-		reference		
ISO 7176-11	1992-05	N	Wheelchairs; part_11: test dummies					
	1		Wheelchairs; part_13: determination of	Y	Y			INL =0.00/10
ISO 7176-13	1989-08	Ν	coefficient of friction of test surfaces	Y	Y	ABNT NBR ISO 7176-13	2009	IN nº 09/13

[Wheelchairs Part_14: Power and control					
			systems for electrically powered wheelchairs					
				Y	Y			IN nº 09/13
100 7470 44	0000 00	N	and scooters Requirements and test			There is no national		
ISO 7176-14	2008-02	N	methods			reference		
			Wheelchairs Part_15: Requirements for					
100 7170 15	1000 11		information disclosure, documentation and					
ISO 7176-15	1996-11	N	labelling					
			Wheelchairs Part_16: Resistance to					
			ignition of upholstered parts Requirements	Y	Y	There is no national		IN nº 09/13
ISO 7176-16	1997-05	N	and test methods			reference		
			Wheelchairs Part_19: Wheeled mobility	Y	Y	There is no national		IN nº 09/13
ISO 7176-19	2008-07	N	devices for use as seats in motor vehicles	•		reference		
			Wheelchairs Part_2: Determination of	Y	Y	There is no national		IN nº 09/13
ISO 7176-2	2001-06	N	dynamic stability of electric wheelchairs	•	•	reference		
			Wheelchairs Part_21: Requirements and					
			test methods for electromagnetic					
			compatibility of electrically powered	Y	Y			IN nº 09/13
			wheelchairs and scooters, and battery			There is no national		
ISO 7176-21	2009-04	N	chargers			reference		
				Y	Y	ABNT NBR ISO 7176-22		IN nº 09/13
ISO 7176-22	2000-05	N	Wheelchairs Part_22: Set-up procedures	•			2009	
			Wheelchairs Part_23: Requirements and					
			test methods for attendant-operated stair-					
ISO 7176-23	2002-07	N	climbing devices					
			Wheelchairs Part_24: Requirements and					
			test methods for user-operated stair-					
ISO 7176-24	2004-10	N	climbing devices					
ISO 7176-26	2007-04	N	Wheelchairs Part_26: Vocabulary					
			Wheelchairs Part_3: Determination of	Y	Y	ABNT NBR ISO 7176-3		IN nº 09/13
ISO 7176-3	2003-04	N	effectiveness of brakes		•		2009	
			Wheelchairs Part_4: Energy consumption					
			of electric wheelchairs and scooters for	Y	Y	There is no national		IN nº 09/13
ISO 7176-4	2008-10	N	determination of theoretical distance range			reference		
1307170-4	2000-10	IN				Telefence		
			Wheelchairs Part_5: Determination of	Y	Y	There is no national		IN nº 09/13
ISO 7176-5	2008-06	Ν	dimensions, mass and manoeuvring space			reference		1111 00/10
10011100	2000 00		Wheelchairs Part_6: Determination of			Telefolie		
			maximum speed, acceleration and	Y	Y	There is no national		IN nº 09/13
ISO 7176-6	2001-10	Ν	deceleration of electric wheelchairs	I	I	reference		INTE 03/13
	2001 10		Wheelchairs Part_7: Measurement of					+
ISO 7176-7	1998-05	Ν	seating and wheel dimensions	Y	Y	ABNT NBR ISO 7176-7	2009	IN nº 09/13
	1000 00		Wheelchairs Part_8: Requirements and					
			test methods for static, impact and fatigue	Y	Y	ABNT NBR ISO 7176-8	2009	IN nº 09/13
ISO 7176-8	1998-07	N	strengths				2000	1111 00/10
100 1110-0	1000-07	IN IN	ouoliguio					

ISO 7176-9 2		N	Wheelchairs Part_9: Climatic tests for electric wheelchairs	Y	Y	There is no national reference	IN nº 09/13
	2009-11					Telefence	
ISO 7193 1	1985-12	Ν	Wheelchairs; Maximum overall dimensions				
			Neurosurgical implants Sterile, single-				
			use hydrocephalus shunts and				
ISO 7197 2	2006-06	Ν	components				
			Neurosurgical implants Sterile, single-				
			use hydrocephalus shunts and				
ISO 7197 Tech 2	2007-07	Ν	components; Technical Corrigendum_1				
			Cardiovascular implants Tubular				
ISO 7198 1	1998-08	Ν	vascular prostheses				
			Cardiovascular implants and artificial				
			organs Blood-gas exchangers				
ISO 7199 2	2009-04	N	(oxygenators)				
			Cardiovascular implants and artificial				
			organs Blood-gas exchangers				
			(oxygenators) Amendment_1:				
			Clarifications for test methodologies,				
ISO 7199 AMD 2	2012-02	N	labelling, and sampling schedule				
			Implants for surgery Partial and total				
			hip joint prostheses Part_1:				
			Classification and designation of				
ISO 7206-1 2	2008-04	N	dimensions				
			Implants for surgery Partial and total				
			hip-joint prostheses Part_10:	Y	Y		
			Determination of resistance to static				
ISO 7206-10 2	2003-12	N	load of modular femoral heads				
			Implants for surgery Partial and total				
			hip joint prostheses Part_2:	Y	Y		
			Articulating surfaces made of metallic,				
ISO 7206-2 2	2011-04	Ν	ceramic and plastics materials				
			Implants for surgery Partial and total			1	
			hip joint prostheses Part_4:				
			Determination of endurance properties				
			and performance of stemmed femoral				
ISO 7206-4 2	2010-06	N	components				

			Implants for surgery; partial and total hip				
			joint prostheses; part_6: determination				
			of endurance properties of head and				
			neck region of stemmed femoral				
ISO 7206-6	1992-03	N	components				
			Implanta far aurgan (Compananta far				
			Implants for surgery Components for				
			partial and total knee joint prostheses Part_1: Classification, definitions and				
ISO 7207-1	2007-02	N	designation of dimensions				
150 7207-1	2007-02	IN	designation of dimensions				
			Implants for surgery Components for				
			partial and total knee joint prostheses	Y	Y		
			Part_2: Articulating surfaces made of				
ISO 7207-2	2011-07	Ν	metal, ceramic and plastics materials				
			Anaesthetic and respiratory equipment	Y	Y		
ISO 7376	2009-08	N	Laryngoscopes for tracheal intubation				
			Medical gas pipeline systems Part_1:				
			Pipeline systems for compressed				
ISO 7396-1	2007-04	Ν	medical gases and vacuum				
100 / 000 1	2007 04						
			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				
ISO 7396-1 A	M2010-01	Ν	the pipeline through flexible hoses				
			Medical gas pipeline systems Part_1:				
			Pipeline systems for compressed	Y	Y		
			medical gases and vacuum;	Ŷ	Ϋ́		
ISO 7396-1 A	M 2010-02	Ν	Amendment_2				
			Medical gas pipeline systems Part_2:				
			Anaesthetic gas scavenging disposal	Y	Y		
ISO 7396-2	2007-04	N	systems				
ISO 7405	2008-12	N	Dentistry Evaluation of biocompatibility of medical devices used in dentistry				
130 7400	2000-12	IN	medical devices used in defilisly		I	1	I

			Copper-bearing contraceptive intrauterine					
ISO 7439	2011-06	Ν	devices Requirements and tests	Y	Y	ABNT NBR ISO 7439	2014	RDC 69/2009
ISO 7488	1991-06	N	Dental amalgamators					
100 / 400	1331-00		Dental materials Determination of colour					
ISO 7491	2000-09	Ν	stability					
ISO 7492	1997-02	N	Dental explorers					
ISO 7493	2006-05	N	Dentistry Operator's stool					
100 / 400	2000 00		Dentistry Dental units Part_1: General					
ISO 7494-1	2011-08	N	requirements and test methods					
	2011.00		Dentistry Dental units Part_2: Water and					
ISO 7494-2	2003-03	Ν	air supply					
ISO 7551	1996-12	N	Dental absorbent points					
			Dental rotary instruments Diamond					
			instruments Part_1: Dimensions,					
ISO 7711-1	1997-02	Ν	requirements, marking and packaging					
			Dental rotary instruments Diamond					
			instruments Part_1: Dimensions,					
			requirements, marking and packaging;					
ISO 7711-1 AM	MD 2009-05	Ν	Amendment_1					
			Dentistry Rotary diamond instruments					
ISO 7711-2	2011-07	Ν	Part_2: Discs					
			Dentistry Diamond rotary instruments					
			Part_3: Grit sizes, designation and colour					
ISO 7711-3	2004-11	N	code					
			Instruments for surgery; Scalpels with					
ISO 7740	1985-12	Ν	detachable blades; Fitting dimensions					
			Instruments for surgery; Scissors and					
			shears; General requirements and test					
ISO 7741	1985-12	Ν	methods					
130 7 741	1905-12	IN	Dental handpieces Part_1: High-speed air					
ISO 7785-1	1997-08	N	turbine handpieces					
100 1100-1	1337-00		Dental handpieces Part_2: Straight and					
ISO 7785-2	1995-08	Ν	geared angle handpieces					
100 1103-2	1333-00		Dental rotary instruments Laboratory					
ISO 7786	2001-04	Ν	abrasive instruments					
100 1100	2001.04		Dental rotary instruments; Cutters; Part 1 :					
ISO 7787-1	1984-12	N	Steel laboratory cutters					
	1004 12		Dental rotary instruments Cutters					
ISO 7787-2	2000-12	Ν	Part 2: Carbide laboratory cutters	Y	NO	ABNT NBR ISO	1999	
	2000 12		Dental rotary instruments; cutters; part_3:		1			1
			carbide laboratory cutters for milling					
ISO 7787-3	1991-12	Ν	machines					
					1	İ		
			Dental rotary instruments Cutters					
ISO 7787-4	2002-03	Ν	Part_4: Miniature carbide laboratory cutters					

			Sterile hypodermic needles for single	Y	Y	ABNT NBR ISO	2010	RDC 05/2011
ISO 7864	1993-05	N	use	I	I.		2010	1100 03/2011
			Dentistry Sterile injection needles for		Y			
ISO 7885	2010-02	N	single use		I			RDC 05/2011
			Sterile hypodermic syringes for single	Y	Y			
ISO 7886-1	1993-10	N	use; part_1: syringes for manual use	Ι	I	ABNT NBR ISO	2003	RDC 03/2011
			Sterile hypodermic syringes for single					
			use Part_1: Syringes for manual use;	Y	Y			
ISO 7886-1 Te	1995-11	N	Technical Corrigendum_1			ABNT NBR ISO	2003	RDC 03/2011
			Sterile hypodermic syringes for single					
			use Part_2: Syringes for use with	Y	Y			
ISO 7886-2	1996-05	Ν	power-driven syringe pumps			ABNT NBR ISO	2003	RDC 03/2011
			Sterile hypodermic syringes for single					
			use Part_3: Auto-disable syringes for					
ISO 7886-3	2005-03	N	fixed-dose immunization					
			Sterile hypodermic syringes for single					
			use Part_4: Syringes with re-use	Y	NO	ABNT NBR ISO	1996	
ISO 7886-4	2006-10	N	prevention feature					
			Optics and optical instruments	Y	NO	ABNT NBR ISO	4007	
ISO 7944	1998-06	N	Reference wavelengths	Ŷ	NO	ABINT INDR ISO	1997	
			Optics and optical instruments					
			Reference wavelengths; Technical					
ISO 7944 Tech	2009-07	Ν	Corrigendum_1					
			Ophthalmic optics Spectacle frames					
			Lists of equivalent terms and					
ISO 7998	2005-10	Ν	vocabulary					
			Mechanical contraceptives Reusable					
			natural and silicone rubber contraceptive					
ISO 8009	2004-10	N	diaphragms Requirements and tests					
			Mechanical contraceptives Reusable					
			natural and silicone rubber contraceptive diaphragms Requirements and tests;					
ISO 8009 AMD 1	2012-02	N	Amendment 1					
ICC COCC AND	2012-02		Small-bore connectors for liquids and					
			gases in healthcare applications					
ISO 80369-1	2010-12	N	Part_1: General requirements					
100 00003-1	2010-12	IN	Medical electrical equipment Part_2-					
			12: Particular requirements for basic					
			safety and essential performance of	Y	Y	There is no notional		IN nº 09/13
ISO 80601-2-1	12011-04	N	critical care ventilators			There is no national		
130 00001-2-1	42011-04	IN				reference		

ISO 80601-2-1:	2011-10	N	Medical electrical equipment Part_2- 12: Particular requirements for basic safety and essential performance of critical care ventilators; Technical Corrigendum_1	Y	Y	There is no national reference		IN nº 09/13
ISO 80601-2-1:	2011-08	N	Medical electrical equipment Part_2- 13: Particular requirements for basic safety and essential performance of an anaesthetic workstation	Y	Y	There is no national reference		IN nº 09/13
ISO 80601-2-5	2011-12	Ν	Medical electrical equipment Part_2- 55: Particular requirements for the basic safety and essential performance of respiratory gas monitors					
ISO 80601-2-56	2009-10	N	Medical electrical equipment Part_2-56: Particular requirements for basic safety and essential performance of clinical thermometers for body temperature measurement	Y	Y	ABNT NBR ISSO/IEC 60601- 2-56	2013	IN nº 09/13
ISO 80601-2-6		N	Medical electrical equipment Part_2- 61: Particular requirements for basic safety and essential performance of pulse oximeter equipment	Y	Y	There is no national reference		IN nº 09/13
ISO 81060-1	2007-12	N	Non-invasive sphygmomanometers Part_1: Requirements and test methods for non-automated measurement type					
ISO 81060-2	2009-05	N	Non-invasive sphygmomanometers Part_2: Clinical validation of automated measurement type Non-invasive sphygmomanometers					
ISO 81060-2 T	2011-02	N	Part_2: Clinical validation of automated measurement type; Technical Corrigendum_1					
ISO 8185	2007-07	N	Respiratory tract humidifiers for medical use Particular requirements for respiratory humidification systems					
ISO 8194	1987-06	N	Radiation protection; Clothing for protection against radioactive contamination; Design, selection, testing and use					

			Acoustics Audiometric test methods					
			Part_1: Pure-tone air and bone					
ISO 8253-1	2010-11	Ν	conduction audiometry					
			Acoustics Audiometric test methods					
			Part_2: Sound field audiometry with					
ISO 8253-2	2009-12	Ν	pure-tone and narrow-band test signals					
			Acoustics Audiometric test methods					
ISO 8253-3	2012-03	N	Part_3: Speech audiometry					
	1001 10	N	Dental equipment Mercury and alloy mixers and dispensers					
ISO 8282	1994-10	N	mixers and dispensers					
			Orthopaedic instruments Drive					
			connections Part_1: Keys for use with					
ISO 8319-1	1996-05	Ν	screws with hexagon socket heads					
			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					
			Oxygen concentrators for medical use	Y	NO	ABNT NBR ISO	1998	
ISO 8359	1996-12	N	Safety requirements	I	NO		1550	
			Injection containers and accessories					
			Part_1: Injection vials made of glass	Y	NO	ABNT NBR ISO	1998	
ISO 8362-1	2009-12	N	tubing					
			Injection containers and accessories					
ISO 8362-2	2008-10	N	Part_2: Closures for injection vials					
			Injection containers and accessories					
			Part_3: Aluminium caps for injection					
ISO 8362-3	2001-12	N	vials					
			Injection containers and accessories					
			Part_4: Injection vials made of moulded					
ISO 8362-4	2011-09	N	glass					
			Injection containers and accessories					
			Part_5: Freeze drying closures for					
ISO 8362-5	2008-10	N	injection vials					

ISO 8362-6	2010-06	N	Injection containers and accessories Part_6: Caps made of aluminium- plastics combinations for injection vials					
130 0302-0	2010-00		Injection containers and accessories Part_7: Injection caps made of					
ISO 8362-7	2006-04	N	aluminium-plastics combinations without overlapping plastics part					
			Optics and optical instruments;					
ISO 8429	1986-09	N	Ophthalmology; Graduated dial scale Infusion equipment for medical use					
ISO 8536-1	2011-09	Ν	Part_1: Infusion glass bottles					
			Infusion equipment for medical use Part_10: Accessories for fluid lines for	Y	Y	ABNT NBR ISO	2004	RDC 04/2011
SO 8536-10	2004-10	N	use with pressure infusion equipment Infusion equipment for medical use					
ISO 8536-11	2004-10	N	Part_11: Infusion filters for use with pressure infusion equipment	Y	Y	ABNT NBR ISO	2004	RDC 04/2011
			Infusion equipment for medical use	Y	NO	ABNT NBR ISO	1996	
SO 8536-12	2007-04	N	Part_12: Check valves	I	NO		1000	
ISO 8536-2	2010-03	N	Infusion equipment for medical use Part_2: Closures for infusion bottles					
			Infusion equipment for medical use					
ISO 8536-3	2009-06	N	Part_3: Aluminium caps for infusion bottles					
30 8530-3	2009-00	IN	Infusion equipment for medical use					
			Part_4: Infusion sets for single use,	Y	Y			
ISO 8536-4	2010-10	N	gravity feed			ABNT NBR ISO	2011	RDC 04/2011
			Infusion equipment for medical use	X	N N			
ISO 8536-5	2004-02	N	Part_5: Burette infusion sets for single use, gravity feed	Y	Y	ABNT NBR ISSO	2012	RDC 04/2011
130 6536-5	2004-02	IN	Infusion equipment for medical use			ADINT INDR 1330	2012	RDC 04/2011
			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-					
0.0526 7	2000.01		plastics combinations for infusion					
ISO 8536-7	2009-01	N	bottles					

			Infusion equipment for medical use					
			Part_8: Infusion equipment for use with	Y	Y	ABNT NBR ISO	2012	RDC 04/2011
ISO 8536-8	2004-08	N	pressure infusion apparatus					
			Infusion equipment for medical use					
			Part_9: Fluid lines for use with pressure	Y	Y	ABNT NBR ISO	2013	RDC 04/2011
ISO 8536-9	2004-10	N	infusion equipment					
			Sterile single-use syringes, with or	Y	Y			
ISO 8537	2007-10	Ν	without needle, for insulin	I	I	ABNT NBR ISO	2012	RDC 03/2011
			Prosthetics and orthotics; limb					
			deficiencies; part_1: method of					
			describing limb deficiencies present at					
ISO 8548-1	1989-08	N	birth					
			Prosthetics and orthotics; limb					
			deficiencies; part_2: method of					
			describing lower limb amputation					
ISO 8548-2	1993-07	N	stumps					
			Prosthetics and orthotics; limb					
			deficiencies; part_3: method of					
			describing upper limb amputation					
ISO 8548-3	1993-07	Ν	stumps					
			Prosthetics and orthotics Limb					
			deficiencies Part_4: Description of					
ISO 8548-4	1998-07	Ν	causal conditions leading to amputation					
			Prosthetics and orthotics Limb					
			deficiencies Part_5: Description of the					
			clinical condition of the person who has					
ISO 8548-5	2003-07	N	had an amputation					
			Prosthetics and orthotics; vocabulary;					
			part_1: general terms for external limb					
ISO 8549-1	1989-07	N	protheses and external orthoses					
			Prosthetics and orthotics; vocabulary;					
ISO 8549-2	1989-07	NI	part_2: terms relating to external limb					
130 0049-2	1909-07	N	prostheses and wearers of these prostheses Prosthetics and orthotics; vocabulary;					
ISO 8549-3	1989-07	Ν	part_3: terms relating to external orthoses					
100 00 00 00 0	1000-07		Prosthetics and orthotics Functional					
			deficiencies Description of the person to					
			be treated with an orthosis, clinical					
			objectives of treatment, and functional					
ISO 8551	2003-08	Ν	requirements of the orthosis					

			Ophthalmic optics Visual acuity			
			testing Standard optotype and its			
ISO 8596	2009-07	N	presentation			
	2000 0.		Optics and optical instruments			
ISO 8598	1996-08	N	Focimeters			
			Optics and optical instruments			
ISO 8598 Tech	n 1998-05	Ν	Focimeters; Technical corrigendum_1			
			Optics and photonics Medical			
			endoscopes and endotherapy devices			
ISO 8600-1	2005-05	N	Part_1: General requirements			
			Optics and optical instruments			
			Medical endoscopes and endoscopic			
			accessories Part_2: Particular			
ISO 8600-2	2002-08	N	requirements for rigid bronchoscopes			
			Optics and optical instruments			
			Medical endoscopes and endoscopic			
			accessories Part_3: Determination of			
			field of view and direction of view of			
ISO 8600-3	1997-07	N	endoscopes with optics			
			Optics and optical instruments			
			Medical endoscopes and endoscopic			
			accessories Part_3: Determination of field of view and direction of view of			
ISO 8600-3 AN	12002 12	N	endoscopes with optics; Amendment_1			
130 8000-3 AN	12003-12	IN	Optics and optical instruments			
			Medical endoscopes and certain			
			accessories Part_4: Determination of			
ISO 8600-4	1997-07	Ν	maximum width of insertion portion			
	1007 07		Optics and photonics Medical			
			endoscopes and endotherapy devices			
			Part_5: Determination of optical			
			resolution of rigid endoscopes with			
ISO 8600-5	2005-03	Ν	optics			
· · · · · ·			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			

		1				
			Implants for surgery; fixation devices for			
ISO 8615	1991-11	Ν	use in the ends of the femur in adults			
130 0015	1991-11	IN				
			Ophthalmic optics Spectacle frames			
100 0004	0011.00	N				
ISO 8624	2011-02	N	Measuring system and terminology			
			O and 's second and 's such as to second			
			Cardiovascular implants and			
			extracorporeal systems			
			Haemodialysers, haemodiafilters,			
ISO 8637	2010-07	N	haemofilters and haemoconcentrators			
			Cardiovascular implants and			
			extracorporeal systems			
			Extracorporeal blood circuit for			
			haemodialysers, haemodiafilters and			
ISO 8638	2010-07	N	haemofilters			
ISO 8669-1	1988-07	N	Urine collection bags; part_1: vocabulary			
			Urine collection bags Part_2:			
ISO 8669-2	1996-12	N	Requirements and test methods			
ISO 8670-1	1988-07	N	Ostomy collection bags; part_1: vocabulary			
100 0010 1	1000 01		Ostomy collection bags, Part_1: voodbulary			
ISO 8670-2	1996-12	Ν	Requirements and test methods			
			Ostomy collection bags Part_3:			
			Determination of odour transmission of			
ISO 8670-3	2000-03	Ν	colostomy and ileostomy bags			
			Implants for surgery; staples with			
			parallel legs for orthopaedic use;			
ISO 8827	1988-10	N	general requirements			
			Implants for surgery; guidance on care			
ISO 8828	1988-10	N	and handling of orthopaedic implants			
			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		Suction catheters for use in the			+
ISO 8836	2007-09	Ν	respiratory tract			
130 0030	2007-09	IN		I		

			Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part_1: Extractables in aqueous			
ISO 8871-1	2003-10	N	autoclavates			
100 0071 1	2000 10		Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part 2: Identification and			
ISO 8871-2	2003-10	Ν	characterization			
100 0071 2	2000 10		Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part_2: Identification and			
ISO 8871-2 A	M 2005-07	N	characterization; Amendment_1			
100 0011 274	2000 01		Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part_3: Determination of released-			
ISO 8871-3	2003-08	N	particle count			
			Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part_4: Biological requirements and test			
ISO 8871-4	2006-06	Ν	methods			
			Elastomeric parts for parenterals and			
			for devices for pharmaceutical use			
			Part_5: Functional requirements and			
ISO 8871-5	2005-08	N	testing			
			Aluminium caps for transfusion, infusion			
			and injection bottles General			
ISO 8872	2003-03	N	requirements and test methods			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_1:			
			Specifications for single-vision and			
ISO 8980-1	2004-02	N	multifocal lenses			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_1:			
			Specifications for single-vision and			
			multifocal lenses; Technical			
ISO 8980-1 Te	e¢2006-08	N	Corrigendum_1			
			Ophthalmic optics Uncut finished			
			spectacle lenses Part_2:			
			Specifications for progressive power			
ISO 8980-2	2004-02	N	lenses			

			Ophthalmic optics Uncut finished					
			spectacle lenses Part_2:					
			Specifications for progressive power					
ISO 8980-2 Te	2006-08	Ν	lenses; Technical Corrigendum_1					
			Ophthalmic optics Uncut finished					
			spectacle lenses Part_3:					
			Transmittance specifications and test					
ISO 8980-3	2003-10	Ν	methods					
			Ophthalmic optics Uncut finished					
			spectacle lenses Part_4:					
			Specifications and test methods for anti-					
ISO 8980-4	2006-08	Ν	reflective coatings					
			Ophthalmic optics Uncut finished					
			spectacle lenses Part_5: Minimum					
			requirements for spectacle lens					
			surfaces claimed to be abrasion-					
ISO 8980-5	2005-08	N	resistant					
			Dentistry Hose connectors for air driven					
ISO 9168	2009-07	Ν	dental handpieces					
			Terminal units for medical gas pipeline					
			systems Part_1: Terminal units for	Y	Y	ABNT ISO/TR	2008	RDC 56 /2001
			use with compressed medical gases	1			2000	100 30 /2001
ISO 9170-1	2008-07	N	and vacuum					
			Terminal units for medical gas pipeline					
			systems Part_2: Terminal units for					
ISO 9170-2	2008-07	N	anaesthetic gas scavenging systems					
			Dentistry Extraction forceps Part_1:					
ISO 9173-1	2006-06	N	General requirements and test methods					
ISO 9173-2	2010-05	N	Dentistry Extraction forceps Part_2: Designation					
130 9173-2	2010-03	11	Injection equipment for medical use					
ISO 9187-1	2010-10	N	Part_1: Ampoules for injectables					
100 9107-1	2010-10		Tart_T. Ampoules for injectables					
			Injection equipment for medical use					
ISO 9187-2	2010-10	Ν	Part_2: One-point-cut (OPC) ampoules					
100 0107 2	2010 10		Implants for surgery; metal bone screws					
			with conical under-surface of head;					
ISO 9268	1988-12	N	dimensions					
100 0200	1000 12		Implants for surgery; metal bone plates;					
			holes and slots corresponding to screws					
ISO 9269	1988-12	N	with conical under-surface					
100 3203	1000-12	IN	พายา ออกเอสา นาเนอา-วินาเสออ					

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	N	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			
	2000					
			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 0000 1	2000 00					
			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			
130 9300-2	2001-04	IN				
			Power-operated lifting platforms for persons			
			with impaired mobility Rules for safety,			
			dimensions and functional operation			
ISO 9386-1	2000-11	N	Part_1: Vertical lifting platforms			
			Power-operated lifting platforms for persons			
			with impaired mobility Rules for safety,			
			dimensions and functional operation			
			Part_2: Powered stairlifts for seated,			
ISO 9386-2	2000-11	N	standing and wheelchair users moving in an inclined plane			
150 9500-2	2000-11		Ophthalmic optics Contact lenses and			
			contact lens care products			
			Determination of biocompatibility by			
ISO 9394	1998-08	N	ocular study with rabbit eyes			
130 9394	1990-00	IN				
			Implants for surgery; non-destructive			
	1002.10	NI NI	testing; liquid penetrant inspection of			
ISO 9583	1993-10	N	metallic surgical implants			

			Implants for surgery; non-destructive					
			testing; radiographic examination of					
ISO 9584	1993-10	Ν	cast metallic surgical implants					
100 0001	1000 10		Implants for surgery; determination of					
			bending strength and stiffness of bone					
ISO 9585	1990-12	Ν	plates					
100 3000	1990-12		Stainless steel needle tubing for					
ISO 9626	1991-09	Ν	manufacture of medical devices	Y	Y	ABNT NBR ISSO	2003	RDC 05/2011
130 9020	1991-09	IN	Stainless steel needle tubing for the			ABINT INBR 1330	2003	RDC 05/2011
				V	V		0000	
	D 0004 00	N	manufacture of medical devices;	Y	Y	ABNT NBR ISSO	2003	RDC 05/2011
ISO 9626 AM	D 2001-06	N	Amendment_1			There is no national		
ISO 9680	2007-06	N	Dentistry Operating lights	Y	Y	reference		IN nº 09/13
ISO 9687	1993-02	N	Dental equipment; graphical symbols			Telefence		
ISO 9693	1999-12	N	Metal-ceramic dental restorative systems					
			Metal-ceramic dental restorative systems;					
ISO 9693 AMD	1 2005-10	Ν	Amendment 1					
			Dentistry Compatibility testing Part_1:					
ISO 9693-1	2012-02	Ν	Metal-ceramic systems					
			Neurosurgical implants Self-closing					
ISO 9713	2002-09	N	intracranial aneurysm clips					
			Orthopaedic drilling instruments; part_1:					
ISO 9714-1	1991-03	N	drill bits, taps and countersink cutters					
			Ophthalmic instruments Trial case					
ISO 9801	2009-12	N	lenses					
			Dental hand instruments Reusable mirrors	N/			0010	
ISO 9873	1998-11	Ν	and handles	Y	NO	ABNT NBR ISO	2013	
			Dental hand instruments Reusable mirrors					
ISO 9873 Tech	ni 2000-06	N	and handles; Technical Corrigendum_1					
			Dentistry Water-based cements Part_1:					
ISO 9917-1	2007-10	N	Powder/liquid acid-base cements					
100 0047 0	0010.01	N	Dentistry Water-based cements Part_2:					
ISO 9917-2	2010-04	N	Resin-modified cements Urine absorbing aids; vocabulary; part_1:					
ISO 9949-1	1993-07	Ν	conditions of urinary incontinence					
130 9949-1	1993-07	IN	Urine absorbing aids; vocabulary; part_2:					
ISO 9949-2	1993-07	Ν	products					
100 0040 2	1000 01		Urine absorbing aids; vocabulary; part_3:					
ISO 9949-3	1993-07	Ν	identification of product types					
ISO 9997	1999-12	N	Dental cartridge syringes					
			Assistive products for persons with					
ISO 9999	2011-07	Ν	disability Classification and terminology					

	1	Electronic Llochth Decord System Eurotional					
0000 44		Electronic Health Record-System Functional					
2009-11	N						
2006-08	N						
2009-07	N						
2009-12	N						
2009-11	N						
		Part_1: Framework for adverse event					
12011-12	N	reporting					
		Health informatics Individual case safety					
2011-12	N						
		Information technology Office equipment					
2008-06	N						
		Requirements and recommendations for					
2011-05	N	interoperability					
2012-05	N						
		Information technology Survey of icons					
		and symbols that provide access to					
		functions and facilities to improve the use of					
		information technology products by the					
2007-07	Ν	elderly and persons with disabilities			<u> </u>		
		Information technology Guidelines for the					
		design of icons and symbols accessible to					
		all users, including the elderly and persons					
2007-06	Ν	with disabilities					
		considerations for people with disabilities					
2009-06	Ν	Part_1: User needs summary					
		Information technology Accessibility					
		considerations for people with disabilities					
2009-06	N	Part_2: Standards inventory					
	2011-12 2008-06 2011-05 2012-05 2007-07 2007-06 2009-06	2006-08 N 2009-07 N 2009-12 N 2009-11 N 20011-12 N 2008-06 N 2011-05 N 2012-05 N 2007-07 N 2007-06 N 2009-06 N	2009-11 N Model, Release_1.1 2006-08 N Reference information model Release_1 2006-08 N Reference information model Release_1 2009-07 N Exchange Standards Health Level Seven Version_2.5 An application protocol for electronic data exchange in healthcare environments 2009-07 N environments 2009-12 N Data Exchange Standards HL7 Clinical Document Architecture, Release_2 2009-11 N services, release_1 Health informatics Common terminology 2009-11 N services, release_1 Health informatics Individual case safety reports (ICSRs) in pharmacovigilance Part_1: Framework for adverse event reports (ICSRs) in pharmacovigilance Part_2: Human pharmacovigilance Part_2: Human pharmacovigilance Part_2: Human pharmacovigilance Part_2: Human pharmacovigilance Part_2: Human pharmacovigilance Part_2: Human pharmacovigilance Part_1: Requirements for ICSR 2011-12 N requirements for ICSR Information technology Office equipment accessibility guidelines for elderly persons and persons with disabilities 2008-06 N and persons with disabilities 2011-05 N information technology Survey of icons and symbols that provide access to functions and facilities to improve the use of information technology	2009-11 N Model, Release_1.1 2006-08 N Health informatics HL_7 version_3 2006-08 N Reference information model Release_1 2006-08 Data Exchange Standards Health Level Seven Version_2.5 An application protocol for electronic data exchange in healthcare 2009-07 N environments 2009-12 N Data Exchange Standards HL7 Clinical 2009-11 N services, release_1 2009-11 N services, release_1 2009-11 N services, release_1 2011-12 N reports (ICSRs) in pharmacovigilance Part_1: Framework for adverse event 2011-12 N reporting 2011-12 N reports (ICSRs) in pharmacovigilance Part_2: Human pharmacoucidal reporting 2011-12 N requirements for ICSR 2011-12 N requirements for ICSR 2011-12 N requirements and recommendations for 2011-12 N requirements and recommendations for 2011-12 N requirements and recommendations for 2008-06 N <td>2009-11 N Model, Release_1.1 2006-08 N Health informaticsHL_7 version_3 2006-08 N Reference information model Release_1 2009-07 N Data Exchange Standards Health Level Seven Version_2.5 An application protocol for electronic data exchange in healthcare </td> <td>2008-11 N Model, Release 1.1 2006-08 N Reference informatios - HL, Z version 3 2006-08 N Reference information modelRelease_1 2008-07 Data Exchange StandardsHealth Level 2009-07 N environments 2009-07 N environments 2009-11 N Document Architecture, Release 2 2009-12 N Document Architecture, Release 2 2009-13 N services, release 1 2009-14 N services, release 1 2009-15 N environments 2009-16 N services, release 1 2011-12 N reporting 2011-12 N reporting 2011-12 N reporting 2011-12 N reguirements for ICSR 2011-15 N information technology, - I</td> <td>2009-11 N Model, Release 1.1 </td>	2009-11 N Model, Release_1.1 2006-08 N Health informaticsHL_7 version_3 2006-08 N Reference information model Release_1 2009-07 N Data Exchange Standards Health Level Seven Version_2.5 An application protocol for electronic data exchange in healthcare	2008-11 N Model, Release 1.1 2006-08 N Reference informatios - HL, Z version 3 2006-08 N Reference information modelRelease_1 2008-07 Data Exchange StandardsHealth Level 2009-07 N environments 2009-07 N environments 2009-11 N Document Architecture, Release 2 2009-12 N Document Architecture, Release 2 2009-13 N services, release 1 2009-14 N services, release 1 2009-15 N environments 2009-16 N services, release 1 2011-12 N reporting 2011-12 N reporting 2011-12 N reporting 2011-12 N reguirements for ICSR 2011-15 N information technology, - I	2009-11 N Model, Release 1.1

			Information technology Accessibility			
			considerations for people with disabilities			
ISO/IEC TR 2913	2009-06	Ν	Part_3: Guidance on user needs mapping			
			Health informatics Point-of-care			
			medical device communication			
ISO/IEEE 1107	2004-12	Ν	Part_10101: Nomenclature			
			Health informatics Point-of-care			
			medical device communication			
ISO/IEEE 1107	2004-12	N	Part_10201: Domain information model			
			Health informatics Personal health device			
			communication Part_10404: Device			
ISO/IEEE 11073-	2010-05	N	specialization Pulse oximeter			
			Health informatics Personal health device			
			communication Part_10407: Device			
ISO/IEEE 11073-	2010-05	Ν	specialization Blood pressure monitor			
			Health informatics Point-of-care			
			medical device communication			
			Part_10408: Device specialization			
ISO/IEEE 1107	2010-05	Ν	Thermometer			
			Health informatics Point-of-care			
			medical device communication			
			Part_10415: Device specialization			
ISO/IEEE 1107	2010-05	Ν	Weighing scale			
			Health informatics Personal health device			
			communication Part_10417: Device			
ISO/IEEE 11073-	2010-05	N	specialization Glucose meter			
			Health informatics Point-of-care			
			medical device communication			
			Part_10471: Device specialization			
ISO/IEEE 1107	2010-05	N	Independant living activity hub			
			Health informatics Point-of care			
			medical device communications			
			Part_20101: Application profiles; Base			
ISO/IEEE 1107	2004-12	N	standard			
			Health informatics Point-of-care			
			medical device communication			
			Part_20601: Application profile			
ISO/IEEE 1107	2010-05	N	Optimized exchange protocol			

			Health informatics Point-of-care			
			medical device communications			
			Part_30200: Transport profile; Cable			
ISO/IEEE 1107	2004-12	N	connected			
130/IEEE 1107	2004-12	IN	Health informatics Point-of-care			
			medical device communications			
			Part_30300: Transport profile; Infrared			
ISO/IEEE 1107	2004-12	N	wireless			
	1000.00		Dental implants; guidelines for developing			
ISO/TR 11175	1993-08	N	dental implants			
	0000 40	N	Health informatics Clinical stakeholder			
ISO/TR 11487	2008-12	N	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	Ν	identifiers and shift marks			
100/11(11040-1	2001-12					
			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			
			Health informatics Information			
			security management for remote			
			maintenance of medical devices and			
			medical information systems Part_1:			
ISO/TR 11633-	2000 11	N				
150/TR 11033-	2009-11	N	Requirements and risk analysis			
			Health informatics Information			
			security management for remote			
			maintenance of medical devices and			
			medical information systems Part_2:			
			Implementation of an information			
ISO/TR 11633-	2009-11	N	security management system (ISMS)			
			Health Informatics Dynamic on-demand			
			virtual private network for health information			
ISO/TR 11636	2009-12	N	infrastructure			
			Guidance on airway management			
ISO/TR 11991	1995-07	Ν	during laser surgery of upper airway			
			Health informatics Guidelines for			
ISO/TR 12309	2009-12	Ν	terminology development organizations			

			Business requirements for health summary			
100 TD 40770 4	0000 00					
ISO/TR 12773-1	2009-06	N	records Part_1: Requirements			
			Business requirements for health summary			
ISO/TR 12773-2	2009-06	N	records Part_2: Environmental scan			
			Medical electrical equipment			
			Deployment, implementation and			
			operational guidelines for indentifying			
			febrile humans using a screening			
ISO/TR 13154	2009-04	N	thermograph			
			Wheelchairs Part_1: Guidelines for the			
			application of the ISO_7176 series on			
ISO/TR 13570-1	2005-04	N	wheelchairs			
ISO/TR 13668	1998-11	N	Digital coding of oral health and care			
	T		Implants for surgery Fundamental			
ISO/TR 14283	2004-07	Ν	principles			
100/11(11200	200101		Health informatics Personal health			
ISO/TR 14292	2012-03	N	records Definition, scope and context			
100/11(11202	2012 00		Dental materials Guidance on testing of			
ISO/TR 14569-1	2007-05	N	wear Part_1: Wear by toothbrushing			
			Medical devices Quality mangement			
			systems Guidance on the application			
ISO/TR 14969	2004 10	N	of ISO_13485: 2003			
130/TK 14909	2004-10	IN	Dentistry Application of OSI clinical			
			codification to the classification and coding			
ISO/TR 15300	2001-05	N	of dental products			
100/111 13300	2001-03		Digital codification of dental laboratory			
ISO/TR 15599	2002-10	N	procedures			
100/11/10000	2002 10		Digital codification of dental laboratory			
ISO/TR 15599 T	2003-10	N	procedures; Technical Corrigendum_1			
	2000 10		Health informatics Interoperability of			
			telehealth systems and networks Part_1:			
ISO/TR 16056-1	2004-07	Ν	Introduction and definitions			
			Health informatics Interoperability of			
			telehealth systems and networks Part_2:			
ISO/TR 16056-2	2004-07	N	Real-time systems			
	T					
			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			
130/1K 10142	2006-01	N	Health informatics Health informatics			
ISO/TR 17119	2005 01	N	profiling framework			
150/1K 1/119	2003-01	N	proming namework		Į	

			Clinical laboratory testing and in vitra		
			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		
			Health informatics Interoperability and		
			compatibility in messaging and		
ISO/TR 18307	2001-12	N	communication standards Key characteristics		
130/11 10307	2001-12	IN	Health informatics Electronic health		
ISO/TR 20514	2005-10	Ν	record Definition, scope and context		
100/11(20011	2000 10				
			Ophthalmic instruments Background		
			for light hazard specification in		
ISO/TR 20824	2007.07	N	ophthalmic instrument standards		
130/18 20024	2007-07	IN	Health informatics Trusted end-to-end		
ISO/TR 21089	2004-06	Ν	information flows		
100/11(21000	200100		Health informatics Security requirements		
			for archiving of electronic health records		
ISO/TR 21548	2010-02	N	Guidelines		
			Health informatics Use of mobile		
			wireless communication and computing		
			technology in healthcare facilities		
			Recommendations for electromagnetic		
			compatibility (management of		
			unintentional electromagnetic		
ISO/TR 21730	2007-02	Ν	interference) with medical devices		
		1	Health informatics Good principles and		
ISO/TR 22221	2006-11	Ν	practices for a clinical data warehouse		
			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	N	persons and persons with disabilities		

T					1	1
			Medical devices utilizing animal tissues			
			and their derivatives Part_4:			
			Principles for elimination and/or			
			inactivation of transmissible spongiform			
			encephalopathy (TSE) agents and			
ISO/TR 22442-2	2010-12	Ν	validation assays for those processes			
			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			
ISO/TR 22676 2	2006-10	Ν	equipment			
			Health informatics Functional			
ISO/TR 22790 2	2007-12	N	characteristics 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			
			Cosmetics Good Manufacturing			
ISO/TR 24475	2010-03	N	Practices General training document			
			Health informatics Business requirements			
			for an international coding system for			
ISO/TR 25257 2	2009-09	N	medicinal products Health informatics Measures for ensuring			
ISO/TR 27809	2007-07	N	patient safety of health software			
130/TR 27009 2	2007-07	IN	Dentistry Guidance on colour			
ISO/TR 28642	2011-07	N	measurement			
100/11(20042 2	2011-07	IN	Ophthalmic optics Spectacle lenses			
			Parameters affecting lens power			
	2007.01	NI	. .			
ISO/TR 28980 2	2007-01	N	measurement			
			Implants for surgery; usage of the terms			
			"valgus" and "varus" in orthopaedic			
ISO/TR 9586	1988-12	N	surgery			
