Status: 2014-07-29								
				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	Mandatory ? Y- fully, P-partial,N- NO	National Reference	Publication date of the national standard	Recognition Number, if available
IEC 60118-0	1983	N Prediant,	Measurement of electroacoustical characteristics	P	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-0 AMD 1	1994-01	N	Hearing aids; part_0: measurement of electroacoustical characteristics; amendment 1	P	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-1	1995-04	N	Hearing aids Part_1: Hearing aids with induction pick-up coil input	Р	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-1 AMD 1	1998-07	N	Hearing aids Part_1: Hearing aids with induction pick-up coil input; Amendment_1	Р	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-1 Edition 3.1	1999-01	N	Hearing aids Part_1: Hearing aids with induction pick-up coil input					
IEC 60118-12	1996-09	N	Hearing aids Part_12: Dimensions of electrical connector systems Electroacoustics Hearing aids Part_13:					
IEC 60118-13	2011-04	N	Electroacousics Hearing aids Part_13. Electromagnetic compatibility (EMC) Hearing aids - Part_14: Specification of a digital interface	Р	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-14	1998-02	N	device Electroacoustics Hearing aids Part_15: Methods for					
IEC 60118-15	2012-02	N	characterising signal processing in hearing aids with a speach-like signal					
IEC 60118-2	1983	N	Hearing aids. Part 2 : Hearing aids with automatic gain control circuits					
IEC 60118-2 AMD 1	1993-02	N	Hearing aids; part_2: hearing aids with automatic gain control circuits; amendment_1					
IEC 60118-2 AMD 2	1997-05	N	Hearing aids Part_2: Hearing aids with automatic gain control circuits; Amendment_2					
IEC 60118-4	2006-10	N	Electroacoustics Hearing aids Part_4: Induction loop systems for hearing aid purposes Magnetic field strength					
IEC 60118-5	1983	N N	Hearing aids. Part 5 : Nipples for insert earphones					
IEC 60118-6	1999-06	N	Hearing aids Part_6: Characteristics of electrical input circuits for hearing aids	Р	N	JIS C 5512:2000	27.03.2000	361 · 362 · 368 · 753
IEC 60118-7	2005-10	N	Electroacoustics Hearing aids Part_7: Measurement of performance characteristics of hearing aids for production, supply and delivery quality assurance purposes					
IEC 60118-8	2005-10	N	Electroacoustics Hearing aids Part_8: Methods of measurement of performance characteristics of hearing aids under simulated in situ working conditions	Р	N	JIS C 5512:2000	27.03.2000	361· 362· 368· 753
IEC 60118-9	1985	N	Hearing aids. Part 9: Methods of measurement of characteristics of hearing aids with bone vibrator output					

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			Electroacoustics - Simulators of human head and ear -					
			Part 4: Occluded-ear simulator for the measurement of					
IEC 60318-4	2010-01	N	earphones coupled to the ear by means of ear inserts					
120 000.0 1	2010 01	.,	Household and similar electrical appliances - Safety -					
			Part_2-52: Particular requirements for oral hygiene					
IEC 60335-2-52	2005-10	N	appliances					
			Household and similar electrical appliances Safety					
			Part_2-52: Particular requirements for oral hygiene					
IEC 60335-2-52 AMD 1	2008-04	N	appliances; Amendment_1					
			Household and similar electrical appliances Safety					
150 00005 0 50 5 W. O.A.	0000 07		Part_2-52: Particular requirements for oral hygiene appliances					
IEC 60335-2-52 Edition 3.1	2008-07	N	- ''					
			Medical electrical equipment X-ray tube					
			assemblies for medical diagnosis Characteristics			JIS Z 4102:2005	2005/3/25	11 · 12 · 14
IEC 60336	2005-04	N	of focal spots	Р	N	JIS Z 4704:2005	2005/3/25	3. 13.
			Medical electrical equipment X-ray tube					
			assemblies for medical diagnosis Characteristics					
IEC 60336 Corrigendum 1	2006-05	N	of focal spots; Corrigendum_1					
3			Determination of the permanent filtration of X-ray					
IEC 60522	2003-12	N	tube assemblies	Р	N	JIS Z 4704:2005	25.03.2005	3. 13.
	2000 12		High-voltage cable plug and socket connections for	· · · · · · · · · · · · · · · · · · ·	.,	0.0 2 11 0 112000	20.00.2000	0 .0
IEC 60526	1978	N	medical X-ray equipment					
120 00020	1370	i v	High-voltage cable plug and socket connections for					
IEC 60526 Corrigendum 1	2010-04	N	medical X-ray equipment					
120 00020 Comgendam	2010 04	i v	Medical electrical equipment - Dose area product					
IEC 60580	2003-09	N	meters					
120 00000	2003 03		Medical electrical equipment - Part 1: General					
			requirements for basic safety and essential					
IEC 60601-1	2005-12	N	performance	Р	N	JIS T 0601-1:2012	01.06.2012	See # 4 (279) on sheet1
	2000 12	.,	Medical electrical equipment - Part 1: General	· · · · · · · · · · · · · · · · · · ·		0.0 1 000 1 1.20 12	0110012012	(270) 511 511 511
			requirements for basic safety and essential					
IEC 60601-1 Corrigendum	2006-12	N	performance; Corrigendum_1	Р	N	JIS T 0601-1:2012	01.06.2012	See # 4 (279) on sheet1
Teo cocor i comgenadii	1 2000 12	.,	Medical electrical equipment - Part 1: General		.,	0.0 1 0001 1.2012	01.00.2012	Coo ii i (270) on onesti
			requirements for basic safety and essential					
IEC 60601-1 Corrigendum	2007-12	N	performance; Corrigendum_2	Р	N	JIS T 0601-1:2012	01.06.2012	See # 4 (279) on sheet1
LC 00001-1 Comgenaum	12007-12	IN	Medical electrical equipment Part_1: General	r	IN	313 1 0001-1.2012	01.00.2012	000 # 4 (213) 011 3110011
			requirements for basic safety and essential					
IEC 60601-1 Interpretation	2009 04	N	performance					
ILC 00001-1 Interpretation	12000-04	IN	Medical electrical equipment Part_1: General					
			requirements for basic safety and essential					
IEC 60601-1 Interpretation	2009-01	N	performance - Interpretation sheet 2					
ILC 00001-1 Interpretation	12003-01	IN	performance interpretation sneet_2					
			Medical electrical equipment Part_1-1: General					
			requirements for safety; Collateral standard: Safety					
IEC 60601-1-1	2000-12	N	requirements for medical electrical systems	Υ	Н	JIS T 0601-1-1:2005	25.03.2005	See # 5 (136) on sheet1
120 0000 1-1-1	2000-12	IN	Medical electrical equipment Part_1-10: General	1	14	010 1 0001-1-1.2000	23.03.2003	555 % 5 (1 55) 511 3116611
			requirements for basic safety and essential					
			performance Collateral Standard: Requirements for the development of physiologic closed-loop					
IEC 60601-1-10	2007-11	N	controllers					
ILC 00001-1-10	2007-11	IN IN	COLITIONE19		1			

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			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					
IEC 60601-1-11	2010-04	N	environment					
			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					
IEC 60601-1-11 Corrigence	2011-04	N	environment					
			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					
IEC 60601-1-11 Technical	2011-04	N	environment; Technical Corrigendum 1					
			Medical electrical equipment Part_1-2: General					
			requirements for basic safety and essential					
			performance Collateral standard:					
			Electromagnetic compatibility Requirements and					
IEC 60601-1-2	2007-03	N	tests	Υ	N	JIS T 0601-1-2:2012	28.03.2012	See # 6 (275) on sheet1
120 00001 12	2007 00	- "	Medical electrical equipment Part_1-2: General		.,	010 1 0001 1 2.2012	20.00.2012	000 # 0 (270) on oneoti
			requirements for basic safety and essential					
			performance Collateral standard:					
			Electromagnetic compatibility Requirements and					
IEC COCO4 4 2 Intermedati	2010 02	NI NI	tests					
IEC 60601-1-2 Interpretati	(2010-03	N	16313					7 · 15 · 16 · 17 · 30 · 397 · 398
			Medical electrical equipment - Part 1-3: General					
			requirements for basic safety and essential performance					· 488 · 569 · 756 · 757 · 482
			Collateral standard: Radiation protection in diagnostic X-					. 1. 2. 3. 4. 5. 6. 8. 9. 10
IEC 60601-1-3	2008-01	N	ray equipment	Υ	N	JIS T 0601-1-3:2012	01.09.2012	· 11· 12· 13· 14· 364
			Medical electrical equipment Part_1: General					
			requirements for safety - 4. Collateral standard:					
IEC 60601-1-4	1996-05	N	Programmable electrical medical systems					
			Medical electrical equipment Part_1-4: General					
			requirements for safety Collateral standard:					
			Programmable electrical medical systems;					
IEC 60601-1-4 AMD 1	1999-10	N	Amendment_1					
			Medical electrical equipment Part_1-4: General					
			requirements for safety Collateral standard:					
IEC 60601-1-4 Edition 1.1	2000-04	N	Programmable electrical medical systems					
			Medical electrical equipment - General					
			requirements for basic safety and essential					
IEC 60601-1-6	2010-01	N	performance - Collateral Standard: Usability					
		+ ''	Medical electrical equipment Part_1-8: General				+	
			requirements for basic safety and essential					
		1	performance Collateral Standard: General					
			i· =					
			requirements, tests and guidance for alarm					
IEC 60604 4 0	2006.40		systems in medical electrical equipment and					
IEC 60601-1-8	2006-10	N	medical electrical systems		1	1		1

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			Madical alastrical agricument - Part 4 Or Canaral					
			Medical electrical equipment Part_1-9: General requirements for basic safety and essential					
			performance Collateral Standard: Requirements					
IEC 60601-1-9	2007-07	N	for environmentally conscious design					
120 00001-1-9	2007-07	IN	Medical electrical equipment Part_2-1: Particular					
			requirements for the basic safety and essential					
			performance of electron accelerators in the range					
IEC 60601-2-1	2009-10	N	1 MeV to 50 MeV					
120 0000121	2000 10	1	Medical electrical equipment; part_2: particular					
			requirements for the safety of nerve and muscle					
IEC 60601-2-10	1987	N	stimulators	Р	N	JIS T 0601-2-10:2005	25.03.2005	788· 123· 124· 141
			Medical electrical equipment - Part 2-10:					
			Particular requirements for the safety of nerve and					
IEC 60601-2-10 AMD 1	2001-09	N	muscle stimulators; Amendment_1	Р	N	JIS T 0601-2-10:2005	25.03.2005	788 · 123 · 124 · 141
			Medical electrical equipment - Part 2-10:					
			Particular requirements for the safety of nerve and					
IEC 60601-2-10 AMD 1 C	d 2002-02	N	muscle stimulators; Amendment_1	Р	N	JIS T 0601-2-10:2005	25.03.2005	788 · 123 · 124 · 141
			Medical electrical equipment - Part 2: Particular					
			requirements for the safety of gamma beam					
IEC 60601-2-11	1997-08	N	therapy equipment	Р	N	JIS Z 4705:2005	25.11.2006	AP7
			Amendment_1 Medical electrical equipment					
			Part_2-11: Particular requirements for the safety of					
IEC 60601-2-11 AMD 1	2004-07	N	gamma beam therapy equipment	Р	N	JIS Z 4705:2005	25.11.2006	AP7
			Medical electrical equipment - Part 2-13:					
			Particular requirements for the safety and essential					
IEC 60601-2-13	2003-05	N	performance of anaesthetic systems					
			Medical electrical equipment - Part 2-13:					
			Particular requirements for the safety and essential					
			performance of anaesthetic systems;					
IEC 60601-2-13 AMD 1	2006-05	N	Amendment_1					
			Medical electrical equipment Part_2-13:					
			Particular requirements for the safety of anaesthetic					
IEC 60601-2-13 Edition 3	.12009-08	N	systems					
			Medical electrical equipment Part_2-16:					
			Particular requirements for basic safety and					
			essential performance of haemodialysis,					
IEC 60601-2-16	2008-04	N	haemodiafiltration and haemofiltration equipment					
			Medical electrical equipment Part_2-16:					
			Particular requirements for basic safety and					
			essential performance of haemodialysis,					
IEC 60601-2-16 Corrigen	d 2008-10	N	haemodiafiltration and haemofiltration equipment					
			Medical electrical equipment Part_2-17:					
			Particular requirements for the safety of					
IEC 60604 0 47	2005 00	N.	automatically-controlled brachytherapy afterloading					
IEC 60601-2-17	2005-09	N	equipment Medical electrical equipment - Part 2-18:					
			Particular requirements for basic safety and					61 · 80 · 55 · 56 · 57 · 58 · 59 ·
IEC 60601-2-18	2009-08	N	essential performance of endoscopic equipment	Υ	N	JIS T 0601-2-18: 2013	2013/9/1	60 · 61 · 62 · 63
IEC 00001-2-18	2009-08	IN	Medical electrical equipment - Part 2-19:	ĭ	IN	010 1 0001-2-10.2013	2013/3/1	00. 01. 07. 03
			Particular requirements for the basic safety and					
IEC 60601-2-19	2009-02	N	essential performance of infant incubators					
ILC 00001-2-19	2009-02	IN	occontial performance of illiant incubators					

		1	Medical electrical equipment Part_2-19:		I		1	
			Particular requirements for the basic safety and					
			essential performance of infant incubators;					
IEC 60601-2-19 Corriger	nd 2012-02	N	Corrigendum 1					
Ö								
			Medical electrical equipment Part_2-2: Particular					
			requirements for the basic safety and essential					
			performance of high frequency surgical equipment					
IEC 60601-2-2	2009-02	N	and high frequency surgical accessories	Р	N	JIS T 0601-2-2: 2014	2014/9/1	150
			Medical electrical equipment - Part 2-20:					
			Particular requirements for the basic safety and					
IEC 60601-2-20	2009-02	N	essential performance of infant transport incubators					
20 00001 2 20	2009-02	IN	Medical electrical equipment Part_2-20:					
			Particular requirements for the basic safety and					
			essential performance of infant transport					
IEC 60601-2-20 Corriger	nd 2012-02	N	incubators; Corrigendum 1					
3			Medical electrical equipment Part_2-21:					
			Particular requirements for the basic safety and					
IEC 60601-2-21	2009-02	N	essential performance of infant radiant warmers	Р	N	JIS T 0601-2-21:2005	25.03.2005	550 119
			Medical electrical equipment Part_2-22:					
			Particular requirements for basic safety and					
			essential performance of surgical, cosmetic,					
IEC 60601-2-22	2007-05	N	therapeutic and diagnostic laser equipment					
			Medical electrical equipment Part_2-23:					
			Particular requirements for the basic safety and					
JEO 00004 0 00	2044.00		essential performance of transcutaneous partial					
IEC 60601-2-23	2011-02	N	pressure monitoring equipment Medical electrical equipment - Part 2-24:		+			
			Particular requirements for the safety of infusion					
IEC 60601-2-24	1998-02	N	pumps and controllers					
120 00001 2 2 1	1000 02		Medical electrical equipment Part_2-25:					
			Particular requirements for basic safety and					
IEC 60601-2-25	2011-10	N	essential performance of electrocardiographs					
			Medical electrical equipment Part_2-26:					
			Particular requirements for the safety of					
IEC 60601-2-26	2003-12	N	electroencephalographs	Р	N	JIS T 1203:1998	30.03.1998	374· 588
			Medical electrical equipment Part_2-27:					
			Particular requirements for the basic safety and					
			essential performance of electrocardiographic					
IEC 60601-2-27	2011-03	N	monitoring equipment	Υ	N			AP14、15
			Medical electrical equipment Part_2-28:					756 1 2 3 4 5 6 8 9
			Particular requirements for basic safety and					· 10· 11· 12· 13· 14· 30· 36
			essential performance of X-ray tube assemblies for					4· 15· 16· 17· 397· 398· 48
IEC 60601-2-28	2010-03	N	medical diagnosis	Υ	N	JIS Z 4751-2-28: 2013	2013/9/1	8 · 569
			Madical destrict environment. Dest. 0.00					
			Medical electrical equipment Part_2-29:					
IEC 60604 2 20	2000.00	l N	Particular requirements for the basic safety and essential performance of radiotherapy simulators	V	N.			A D0
IEC 60601-2-29	2008-06	N	essential performance of faulotherapy simulators	Υ	N			AP8

	1	1	Medical electrical equipment; part_2: particular		1			
			requirements for the safety of short-wave therapy					
IEC 60601-2-3	1991-06	N	equipment	Р	N	JIS T 0601-2-3:2005	25.03.2005	335
			Medical electrical equipment - Part 2: Particular					
			requirements for the safety of short-wave therapy					
IEC 60601-2-3 AMD 1	1998-09	N	equipment; Amendment_1	Р	N	JIS T 0601-2-3:2005	25.03.2005	335
			Medical electrical equipment Part_2-31:					
			Particular requirements for basic safety and					
			essential performance of external cardiac					
IEC 60601-2-31	2008-03	N	pacemakers with internal power source					
			Medical electrical equipment Part_2-31:					
			Particular requirements for basic safety and					
			essential performance of external cardiac					
IEC 60601-2-31 AMD 1	2011-06	N	pacemakers with internal power source					
			Medical electrical equipment Part_2-31:					
			Particular requirements for basic safety and					
			essential performance of external cardiac					
IEC 60601-2-31 Edition 2	.12011-09	N	pacemakers with internal power source					
150 00004 0 00	4004.00		Medical electrical equipment; part_2: particular requirements for the safety of X-ray equipment	-				0 44 40 40 44
IEC 60601-2-32	1994-03	N	Medical electrical equipment - Part 2-33: Particular	Р	N	JIS Z 4703:1995	01.03.1995	. 3. 11. 12. 13. 14.
			requirements for the basic safety and essential					
			performance of magnetic resonance equipment for					
IEC 60601-2-33	2010-03	N	medical diagnosis	Υ	N	JIS Z 4951:2012	01.06.2012	372· 27· 827
			Medical electrical equipment Part_2-33:					
			Particular requirements for the basic safety and					
			essential performance of magnetic resonance					
IEC 60601-2-33 Corrigendu	m 2012-03	N	equipment for medical diagnosis	Υ	N	JIS Z 4951:2012	01.06.2012	372· 27· 827
			Medical electrical equipment Part_2-34:					
			Particular requirements for the basic safety and					
			essential performance of invasive blood pressure					
IEC 60601-2-34	2011-05	N	monitoring equipment	Υ	N	JIS T 0601-2-34:2005	25.03.2005	599 601
			Medical electrical equipment Part_2: Particular					
			requirements for the safety of equipment for					
IEC 60601-2-36	1997-03	N	extracorporeally induced lithotripsy					
			Medical electrical equipment Part_2-37:					
			Particular requirements for the basic safety and					
			essential performance of ultrasonic medical					26 · 57 · 58 · 783 · 20 · 21 · 22
IEC 60601-2-37	2007-08	N	diagnostic and monitoring equipment	Υ	N	JIS T 0601-2-37: 2013	2013/9/1	· 23· 24· 25
			Medical electrical equipment Part_2-39:					
			Particular requirements for basic safety and					
			essential performance of peritoneal dialysis					
IEC 60601-2-39	2007-11	N	equipment	Р	N	JIS T 0601-2-39:2013	01.03.2013	AP35
			Medical electrical equipment Part_2-4: Particular					
.=0		1	requirements for basic safety and essential					
IEC 60601-2-4	2010-12	N	performance of cardiac defibrillators		ļ			
			Madical alastrical assistant Dark 0.40					
			Medical electrical equipment Part_2-40: Particular requirements for the safety of					
JEC 60604 2 40	1000.00		electromyographs and evoked response equipment	Б		IIO T 0004 0 40 0005	0F 00 000F	274. 500
IEC 60601-2-40	1998-02	N	electromyographis and evoked response equipment	Р	N	JIS T 0601-2-40:2005	25.03.2005	374· 599

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			Medical electrical equipment Part_2-41:					
			Particular requirements for basic safety and					
			essential performance of surgical luminaires and					
IEC 60601-2-41	2009-08	N	luminaires for diagnosis					
			Medical electrical equipment Part_2-43:					
			Particular requirements for basic safety and					
			essential performance of X-ray equipment for					
IEC 60601-2-43	2010-03	N	interventional procedures	Υ	N	JIS_Z_4751-2-43:2012	01.10.2012	6 · 364
			Medical electrical equipment Part_2-44:					
			Particular requirements for the basic safety and					
			essential performance of X-ray equipment for					17 · 398 · 488 · 569 · 15 · 364 ·
IEC 60601-2-44	2009-02	N	computed tomography	Υ	N	JIS Z 4751-2-44: 2012	2012/10/1	397
			Medical electrical equipment Part_2-44:					
			Particular requirements for the basic safety and					
			essential performance of X-ray equipment for					
IEC 60601-2-44 Corrige	end 2010-05	N	computed tomography					
<u> </u>			Medical electrical equipment Part_2-45:					
			Particular requirements for the basic safety and					
			essential performance of mammographic X-ray					
			equipment and mammographic stereotactic					
IEC 60601-2-45	2011-02	N	devices	Υ	N	JIS Z 4751-2-45: 2013	2013/9/1	3.7
			Medical electrical equipment Part_2-46:					
			Particular requirements for the basic safety and					
IEC 60601-2-46	2010-12	N	essential performance of operating tables					
			Medical electrical equipment Part_2-47:					
			Particular requirements for the basic safety and					
			essential performance of ambulatory					
IEC 60601-2-47	2012-02	N	electrocardiographic systems					
			Medical electrical equipment Part_2-49:					
			Particular requirements for the basic safety and					
			essential performance of multifunction patient					
IEC 60601-2-49	2011-02	N	monitoring equipment					
			Medical electrical equipment Part_2-5: Particular					
			requirements for basic safety and essential					
IEC 60601-2-5	2009-07	N	performance of ultrasonic physiotherapy equipment	Υ	N	JIS T 0601-2-5:2005	25.03.2005	126-127
			Medical electrical equipment Part_2-50:					
			Particular requirements for the basic safety and					
			essential performance of infant phototherapy					
IEC 60601-2-50	2009-03	N	equipment					
			Medical electrical equipment Part_2-50:					
			Particular requirements for the basic safety and					
			essential performance of infant phototherapy					
IEC 60601-2-50 Corrige	end 2010-08	N	equipment					
			Medical electrical equipment Part_2-52:					
			Particular requirements for the basic safety and					
IEC 60601-2-52	2009-12	N	essential performance of medical beds					
			Medical electrical equipment Part_2-52:					
			Particular requirements for the basic safety and					
IEC 60601-2-52 Corrige	end 2010-09	N	essential performance of medical beds					

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			Medical electrical equipment Part_2-52: Particular requirements for the basic safety and					
			essential performance of medical beds; Technical					
IEC 60601-2-52 Technical	2010-09	N	Corrigendum_1					
120 00001 2 02 10011110a1	2010 00	1	comgondam_1					
			IEC_60601-2-54, Ed1: Medical electrical					
			equipment Part_2-54: Particular requirements for					
			the basic safety and essential performance of X-ray					4· 5· 8· 9· 10· 1· 2· 28· 29
IEC 60601-2-54	2009-06	N	equipment for radiography and radioscopy	Р	N	JIS Z 4751-2-54:2012	01.10.2012	· 482
			Medical electrical equipment Part_2-54:					
			Particular requirements for the basic safety and					
			essential performance of X-ray equipment for					4· 5· 8· 9· 10· 1· 2· 28· 29
IEC 60601-2-54 Corrigend	2010-03	N	radiography and radioscopy	Р	N	JIS Z 4751-2-54:2012	01.10.2012	· 482
			Medical electrical equipment Part_2-54:					
			Particular requirements for the basic safety and					
150 00004 0 54 0	0044.00		essential performance of X-ray equipment for					
IEC 60601-2-54 Corrigend	2011-06	N	radiography and radioscopy					
			Medical electrical equipment Part_2-57: Particular requirements for the basic safety and					
			essential performance of non-laser light source					
			equipment intended for therapeutic, diagnostic,					
IEC 60601-2-57	2011-01	N	monitoring and cosmetic/aesthetic use					
120 00001 2 37	2011 01	111	Medical electrical equipment. Part 2: Particular					
			requirements for the safety of microwave therapy					
IEC 60601-2-6	1984	N	equipment	Р	N	JIS T 0601-2-6:2005	25.03.2005	125
					-			-
			Medical electrical equipment Part_2-7: Particular					
			requirements for the safety of high-voltage					
IEC 60601-2-7	1998-02	N	generators of diagnostic X-ray generators	Υ	N	JIS Z 4751-2-7:2008	25.11.2008	· 3· · 11· 12· 13· 14· AP 7
			Medical electrical equipment Part 2-8: Particular					
			requirements for the basic safety and essential					
			performance of therapeutic X-ray equipment					
IEC 60601-2-8	2010-11	N	operating in the range 10_kV to 1_MV					
			Medical electrical equipment Part_2: Particular					
			requirements for the safety of therapeutic X-ray					
IEC 60601-2-8 AMD 1	1997-08	N	equipment in the range 10_kV to 1_MV; Amendment 1					
IEC 60001-2-8 AIVID 1	1997-08	IN	Amenament_1					
			Medical electrical equipment - Part 2-8: Particular					
			requirements for the safety of therapeutic X-ray					
IEC 60601-2-8 Edition 1.1	1999-04	N	equipment 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					
IEC 60601-3-1	1996-07	N	monitoring equipment					
			Electrical and loading characteristics of X-ray tube			JIS Z 4102:2005	2005/3/25	11. 12. 14
IEC 60613	2010-01	N	assemblies for medical diagnosis	Р	N	JIS Z 4704:2005	2005/3/25	1. 3. 13
			Diagnostic X-ray imaging equipment					
			Characteristics of general purpose and					
IEC 60627	2001-08	N	mammographic anti-scatter grids					

	1		Electroacoustics Audiometric equipment					
IEC 60645-1	2012-02	N	Part_1: Equipment for pure-tone audiometry	Р	N	JIS T 1201-1:2011	29.07.2011	49 · 50 · 52
120 00040-1	2012-02	IN	Audiometers; part_2: equipment for speech	Г	IN	010 1 1201-1.2011	23.01.2011	70 30 32
150 00045 0	1993-11	N	audiometry	Р	N	110 T 4004 0 0000	04.00.0000	49 · 52
IEC 60645-2	1993-11	IN	-	٢	IN	JIS T 1201-2:2000	01.08.2000	49. 32
150 00045 0	0007.00	N.	Electroacoustics Audiometric equipment					
IEC 60645-3	2007-03	N	Part_3: Test signals of short duration					
			Electroacoustics Audiometric equipment					
IEC COCAE E	2004 44	N	Part_5: Instruments for the measurement of aural					
IEC 60645-5	2004-11	IN IN	acoustic impedance/admittance Electroacoustics Audiometric equipment		+			
			Part_6: Instruments for the measurement of					
IEC 60645-6	2009-04	N	otoacoustic emissions					
IEC 60645-6	2009-04	IN	Electroacoustics - Audiometric equipment -					
			Part 7: Instruments for the measurement of					
IEC 60645-7	2009-04	N	_					
ILC 00043-7	2009-04	IN	auditory brainstem responses Medical electrical equipment Characteristics and					
			test conditions of radionuclide imaging devices					
IEC 60789	2005-10	N	Anger type gamma cameras					
IEC 60789	2005-10	IN	Anger type gamma cameras					
			Medical electrical equipment Characteristics and					
			test conditions of radionuclide imaging devices					
IEC 60789 Corrigendum 1	2009-10	N	Anger type gamma cameras; Corrigendum_1					
LC 00709 Conigendani i	2009-10	in in	Determination of the maximum symmetrical					
			radiation field from a rotating anode X-ray tube for					
IEC 60806	1984	N	medical diagnosis					
120 00000	1304		Medical electrical equipment Medical electron					
			accelerators Functional performance					
IEC 60976	2007-10	N	characteristics	Р	N	JIS Z 4714:2001	01.06.2001	AP7
			Safety requirements for electrical equipment for					
			measurement, control and laboratory use Part_2-					
			040: Particular requirements for sterilizers and					
IEC 61010-2-040	2005-04	N	washer-disinfectors used to treat medical materials					
	1		Safety requirements for electrical equipment for					
			measurement, control and laboratory use Part_2-					
			101: Particular requirements for in vitro diagnostic					
IEC 61010-2-101	2002-01	N	(IVD) medical equipment		<u> </u>			
			Standard means for the reporting of the acoustic					
IEC 61157	2007-08	N	output of medical diagnostic ultrasonic equipment					
			Standard means for the reporting of the acoustic					
			output of medical diagnostic ultrasonic equipment;					
IEC 61157 Corrigendum 1	2008-08	N	Corrigendum_1					
			Radiotherapy simulators; functional performance					
IEC 61168	1993-12	N	characteristics	Υ	N			AP8
			Ultrasonics; dental descaler systems; measurement and					
IEC 61205	1993-12	N	declaration of the output characteristics					
			Radiotherapy equipment coordinates, movements					
IEC 61217	2011-12	N	and scales					

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			Evaluation and routine testing in medical imaging					
			departments Part_2-6: Constancy tests					
			Imaging performance of computed tomography X-					
IEC 61223-2-6	2006-11	N	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: Acceptance tests Imaging					
IEC 61223-3-4	2000-03	N	performance of dental X-ray equipment	Υ	N	JIS Z 4752-3-4:2005	25.03.2005	11 · 12 · 13 · 14 · 757
120 01225-5-4	2000-03	- 14	Evaluation and routine testing in medical imaging		- 11	010 2 47 32-3-4.2003	25.05.2005	11 12 10 14 707
			departments Part_3-5: Acceptance tests					
			Imaging performance of computed tomography X-					
IEC 61223-3-5	2004-08	N	ray equipment	Υ	N	JIS Z 4752-3-5:2008	25.06.2008	16
10 01223-3-3	2004-00	IN	Evaluation and routine testing in medical imaging	'	IN	JIS Z 47 52-3-3.2008	25.00.2006	10
			departments Part_3-5: Acceptance tests					
I			Imaging performance of computed tomography X-					
IEC 61223-3-5 Corrige	ndu 2006 03	N	ray equipment; Corrigendum_1					
ILC 01223-3-3 Conige	11du 2000-03	IN						
IFO 040F0 Edition 4.4	2002.02	NI	Electroacoustics Specifications for personal					
IEC 61252 Edition 1.1	2002-03	N	sound exposure meters Medical electrical equipment - Characteristics of					
IEC 61262-1	1994-07	N	electro-optical X-ray image intensifiers Part_1: Determination of the entrance field size					
IEC 01202-1	1994-07	IN	Medical electrical equipment Characteristics of					
			electro-optical X-ray image intensifiers Part_2:					
IEC 61262-2	1994-07	N	Determination of the conversion factor					
ILG 01202-2	1994-07	IN	Medical electrical equipment Characteristics of					
			electro-optical X-ray image intensifiers - Part 3:					
			Determination of the luminance distribution and					
IEC 61262-3	1994-07	N	luminance non-uniformity					
1202 0	1334 01		Medical electrical equipment Characteristics of					
			electro-optical X-ray image intensifiers Part_4:					
IEC 61262-4	1994-07	N	Determination of the image distortion					
120 01202 1	100101	.,	Botomination of the image dictoration					
			Medical electrical equipment - Characteristics of					
			electro-optical X-ray image intensifiers Part_5:					
IEC 61262-5	1994-07	N	Determination of the detective quantum efficiency					
			Medical electrical equipment Characteristics of					
			electro-optical X-ray image intensifiers Part_6:					
			Determination of the contrast ratio and veiling glare					
IEC 61262-6	1994-07	N	index					
			Medical electrical equipment Characteristics of					
			electro-optical X-ray image intensifiers Part-7:					
IEC 61262-7	1995-09	N	Determination of the modulation transfer function					
			Ultrasonics Hand-held probe Doppler foetal					
			heartbeat detectors Performance requirements					
IEC 61266	1994-12	N	and methods of measurement and reporting	Р	N	JIS T 1506:2005	25.03.2005	39
			Medical diagnostic X-ray equipment Radiation					
			conditions for use in the determination of					
IEC 61267	2005-11	N	characetristics					

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		1	Medical electrical equipment Radionuclide			
			calibrators Particular methods for describing			
IEC 61303	1994-09	N	performance			
			Electrical equipment for measurement, control and			
			laboratory use, control and laboratory use EMC			
			requirements Part_2-6: Particular requirements			
IEC 61326-2-6	2005-12	N	In-vitro diagnostic (IVD) medical equipment			
			Electrical equipment for measurement, control and			
			laboratory use, control and laboratory use EMC			
			requirements Part_2-6: Particular requirements			
			In-vitro diagnostic (IVD) medical equipment;			
IEC 61326-2-6 Corrigendo	u 2007-09	N	Corrigendum_1			
			Protective devices against diagnostic medical X-			
			radiation Part_1: Determination of attenuation			
IEC 61331-1	1994-10	N	properties of materials			
			Protective devices against diagnostic medical X-			
IEC 61331-2	1994-10	N	radiation Part_2: Protective glass plates			
			Protective devices against diagnostic medical X-			
.= 0 0 100 1			radiation Part_3: Protective clothing and protective devices for gonads			
IEC 61331-3	1998-11	N				
			Ultrasonics Pulse echo scanners Part_1:			
			Techniques for calibrating spatial measurement			
			systems and measurement of system point-spread			
IEC 61391-1	2006-07	N	function response			
			Ultrasonics Pulse-echo scanners Part_2:			
			Measurement of maximum depth of penetration			
IEC 61391-2	2010-01	N	and local dynamic range			
			Electroacoustics Equipment for the measurement of			
IEC 61669	2001-01	N	real-ear acoustical characteristics of hearing aids			
			Medical electrical equipment Dosimeters with			
			ionization chambers and/or semi-conductor			
			detectors as used in X-ray diagnostic imaging;			
IEC 61674 AMD 1	2002-06	N	Amendment_1			
			Radionuclide imaging devices Characteristics			
			and test conditions Part_1: Positron emission			
IEC 61675-1	1998-02	N	tomographs			
			Radionuclide imaging devices Characteristics			
			and test conditions Part_1: Positron emission			
IEC 61675-1 AMD 1	2008-04	N	tomographs; Amendment_1			
			Radionuclide imaging devices Characteristics			
			and test conditions Part_1: Positron emission			
IEC 61675-1 Edition 1.1	2008-06	N	tomographs			
			Radionuclide imaging devices Characteristics	 		
		1	and test conditions Part_2: Single photon			
IEC 61675-2	1998-01	N	emission computed tomographs			
			Radionuclide imaging devices Characteristics			
		1	and test conditions Part_2: Single photon			
IEC 61675-2 AMD 1	2004-12	N	emission computed tomographs; Amendment_1			
			Radionuclide imaging devices Characteristics			
		1	and test conditions Part_2: Single photon			
IEC 61675-2 Edition 1.1	2005-02	N	emission computed tomographs			
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			Radionuclide imaging devices Characteristics			
			and test conditions Part_3: Gamma camera			
IEC 61675-3	1998-02	N	based wholebody imaging systems			
			Medical electrical equipment Dosimetric			
			instruments used for non-invasive measurement of			
IEC 61676	2002-09	N	X-ray tube voltage in diagnostic radiology			
			Medical electrical equipment Dosimetric			
			instruments used for non-invasive measurement of			
			x-ray tube voltage in diagnostic radiology;			
IEC 61676 AMD 1	2008-11	N	Amendment_1			
			Medical electrical equipment Dosimetric			
			instruments used for non-invasive measurement of			
IEC 61676 Edition 1.1	2009-01	N	X-ray tube voltage in diagnostic radiology			
			Medical electrical equipment Dosimetric instruments used for non-invasive measurement of X-ray tube voltage			
IEC 61685	2002-09	N	in diagnostic radiology			
ILC 01003	2002-09	IN				
			Ultrasonics Physiotherapy systems Field specifications and methods of measurement in the			
IEC 61689	2007-08	N	l ·			
IEC 0 1009	2007-08	IN	frequency range 0,5_MHz to 5_MHz Ultrasonics Pressure pulse lithotripters			
IEC 64046	1998-04	N	Characteristics of fields			
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			Ultrasonics - Surgical systems - Measurement and			
IEC 61847	1998-01	N	declaration of the basic output characteristics			
ILC 01047	1990-01	IN	Medical electrical equipment Requirements for			
			the safety of radiotherapy treatment planning			
IEC 62083	2009-09	N	systems			
ILC 02003	2009-09	IN	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 devices Part_1:			
IEC 62220-1	2003-10	N	Determination of the 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			
			Medical electrical equipment Safety of			
IEC 62274	2005-05	N	radiotheraphy record and verify systems			
			Medical device software Software life cycle			
IEC 62304	2006-05	N	processes			
			Medical electrical equipment Recurrent test and			
IEC 62353	2007-05	N	test after repair of medical electrical equipment			
			Ultrasonics Field characterization Test methods			
			for the determination of thermal and mechanical			
			indices related to medical diagnostic ultrasonic			
IEC 62359	2010-10	N	fields			

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			Ultrasonics Field characterization Test methods					
			for the determination of thermal and mechanical					
			indices related to medical diagnostic ultrasonic					
IEC 62359 Corrigendum 1	2011-03	N	fields					
			Medical devices Application of usability					
IEC 62366	2007-10	N	engineering to medical devices					
			Magnetic resonance equipment for medical imaging					
			Part_1: Determination of essential image quality					
IEC 62464-1	2007-01	N	parameters	Υ	N	JIS Z 4952:2012	01.06.2012	27· 372· 827
			Magnetic resonance equipment for medical imaging					
IEC 62464-2	2010-11	N	Part_2: Classification criteria for pulse sequences					
			Electroacoustics Audio-frequency induction loop					
			systems for assisted hearing Part_1: Methods of measuring and specifying the performance of system					
IEC 00400 4	2010-01	N.	components					
IEC 62489-1	2010-01	N	Electroacoustics - Audio-frequency induction loop					
			systems for assisted hearing Part_2: Methods of					
			calculating and measuring the low-frequency magnetic					
			field emissions from the loop for assessing conformity					
IEC 62489-2	2011-01	N	with guidelines on limits for human exposure					
			Medical electrical equipment Exposure index of					
			digital X-ray imaging systems Part_1: Definition					
IEC 62494-1	2008-08	N	and requirements of general radiography					
120 02494-1	2000-00	IN	Medical electrical equipment - Medical image					
IEC 62563-1	2009-12	N	display systems Part_1: Evaluation methods					
ILC 02303-1	2009-12	IN	Application of risk management for IT-networks					
			incorporating medical devices - Part 1: Roles,					
IEC 80001-1	2010-10	N	responsibilities and activities					
IEC 8000 1-1	2010-10	N	Medical electrical equipment Part_2-30:					
			Particular requirements for basic safety and					
IEO 00004 0 00	0000 04	N	essential performance of automated non-invasive					
IEC 80601-2-30	2009-01	IN .	sphygnomanometers					
			Medical electrical equipment Part_2-30:					
			Particular requirements for basic safety and					
			essential performance of automated non-invasive					
IEC 80601-2-30 Corrigend	2010-01	N	sphygnomanometers; Corrigendum_1					
			Medical electrical equipment Part_2-35:					
			Particular requirements for the basic safety and					
			essential performance of heating devices using					
			blankets, pads and mattresses and intended for					
IEC 80601-2-35	2009-10	N	heating in medical use					
			Medical electrical equipment Part_2-35:					
	1		Particular requirements for the basic safety and					
			essential performance of heating devices using					
			blankets, pads and mattresses and intended for					
IEC 80601-2-35 Corrigend	2012-03	N	heating in medical use					
	1		Medical electrical equipment Part_2-58:					
	1		Particular requirements for basic safety and					
			essential performance of lens removal devices and					
IEC 80601-2-58	2008-10	N	vitrectomy devices for ophthalmic surgery					

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			Medical electrical equipment Part_2-59:				
			Particular requirements for basic safety and				
IEC 80601-2-59	2008-10	N	essential performance of screening thermographs for human febrile temperature screening				
ILC 00001-2-39	2000-10	IN .	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 Corri	gend 2009-04	N	Corrigendum 1				
			Medical electrical equipment Part_2-60: Particular				
IEO 00004 0 00	2012-02		requirements for basic safety and essential performance of dental equipment				
IEC 80601-2-60	2012-02	N					
IEC/TR 60788	2004-02	N	Medical electrical equipment Glossary of defined terms				
IEC/1K 00/00	2004-02	IN	Safety of laser products Part_8: Guidelines for the safe				
IEC/TR 60825-8	2006-12	N	use of laser beams on humans				
			Methods of measuring the performance of				
IEC/TR 60854	1986	N	ultrasonic pulse-echo diagnostic equipment				
			Graphical symbols for electrical equipment in				
IEC/TR 60878	2003-07	N	medical practice				
			Guidelines for administrative, medical, and nursing				
JEO/TD 00000	0000 00	N	staff concerned with the safe use of medical				
IEC/TR 60930	2008-09	N	electrical equipment and medical electrical systems				
			Medical electrical equipment Medical electron accelerators Guidelines for functional				
IEC/TR 60977	2008-07	N	performance characteristics				
120/11/ 003/1	2000 07	1,	periormance characteristics				
			Guidelines for the development and use of medical				
IEC/TR 61258	2008-08	N	electrical equipment educational materials				
			High frequency surgical equipment Operation and				
IEC/TR 61289	2011-11	N	maintenance				
			Nuclear medicine instrumentation Routine tests				
.=0.=B.0			Part_2: Scintillation cameras and single photon				
IEC/TR 61948-2	2001-02	N	emission computed tomography imaging				
IEC/TR 61948-3	2005-07	N	Nuclear medicine instrumentation Routine tests Part_3: Positron emission tomographs				
IEC/TR 01940-3	2005-07	IN	Nuclear medicine instrumentation - Routine tests -				
IEC/TR 61948-4	2006-11	N	Part 4: Radionuclide calibrators				
120/11/010101	2000 11	.,	Medical electrical equipment Guidelines for				
IEC/TR 62266	2002-03	N	implementation of DICOM in radiotherapy				
			Considerations of unaddressed safety aspects in				
			the second edition of IEC_60601-1 and proposals				
IEC/TR 62296	2009-01	N	for new requirements				
LEG/ED 00-1-			Mapping between the clauses of the third edition of				
IEC/TR 62348	2006-05	N	IEC_60601-1 and the 1988 edition as amended				
IEC/TD coops	2000 40	N.	General testing procedures for medical electrical				
IEC/TR 62354	2009-10	N	equipment				
			Requirements for measurement standards for high				
IEC/TR 62649	2010-04	N	intensity therapeutic ultrasound (HITU) devices				
120,111 02040	12010 07	1.4	interiory therapeutic diffasound (11110) devices	l	1		

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			Audio, video and multimedia systems and equipment Activities and considerations related to accessibility and					
IEC/TR 62678	2010-10	N	usability					
IEC/1K 02076	2010-10	IN	Medical device software Part_1: Guidance on the					
IEC/TR 80002-1	2009-09	N	application of ISO 14971 to medical device software					
120/11/ 00002 1	2000 00	.,	Radiotherapy simulators; guidelines for functional					
IEC/TR2 61170	1993-12	N	performance characteristics					
ILC/TRZ 01170	1993-12	IN	Evaluation and routine testing in medical imaging					
IEC/TR2 61223-1	1993-07	N	departments; part_1: general aspects					
ILO/11\Z 01ZZJ-1	1995-01	IN	Ultrasonics - Real-time pulse-echo systems - Test					
			procedures to determine the performance					
IEC/TR2 61390	1996-07	N	specifications					
ILO/11\2 01330	1990-01	IN	Fundamental aspects of safety standards for					
IEC/TR3 60513	1994-01	N	medical electrical equipment					
120/11/0 00010	1334 01	11	Cardiac defibrillators; cardiac defibrillators-					
IEC/TR3 61288-1	1993-10	N	monitors; part 1: operation					
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IEC/TR3 61288-2	1993-10	N	monitors; part 2: maintenance					
120/11/00/1200/2	1333 10	11	Medical electrical equipment Digital imaging and					
			communications in medicine (DICOM)					
IEC/TR3 61852	1998-04	N	Radiotherapy objects					
IEC/TR3 61859	1997-04	N	Guidelines for radiotherapy treatment rooms design					
120/1103 01039	1997-04	IN .	Calabinite in radioantapy a california accign					
			Medical suction equipment Part_1: Electrically					
ISO 10079-1	1999-08	N	powered suction equipment Safety requirements					
130 10079-1	1999-00	IN	Medical suction equipment - Part 2: Manually		+			
ISO 10079-2	1999-08	N	powered suction equipment	Р	N	JIS T 7208-2:2005	25.03.2005	106
130 10079-2	1999-00	IN	Medical suction equipment Part_3: Suction	г	IN	313 1 7208-2.2003	23.03.2003	100
			equipment powered from a vacuum or pressure					
ISO 10079-3	1999-08	N	source					
130 10079-3	1999-00	IN	Oxygen concentrator supply systems for use with					
ISO 10083	2006-07	N	medical gas pipeline systems					
100 10003	2000-07	IN	Dentistry Soft lining materials for removable dentures					
ISO 10139-1	2005-02	N	Part_1: Materials for short-term use					
			Dentistry Soft lining materials for removable dentures					
			Part_1: Materials for short-term use; Technical					
ISO 10139-1 Technical C	Corrig 2006-03	N	Corrigendum_1					
			Dentistry Soft lining materials for removable dentures					
ISO 10139-2	2009-08	N	Part_2: Materials for long-term use	Р	N	JIS T 6520: 2012	2012/7/1	245
			Health informatics Messages and communication					
ISO 10159	2011-12	N	Web access reference manifest					
ISO 10271	2011-08	N	Dentistry Corrosion test methods for metallic materials	Υ	N			AP36
			Single-use sterile rubber surgical gloves					
ISO 10282	2002-09	N	Specification	Р	N	JIS T 9107:2005	25.03.2005	328 · 399 · 400
			Single-use sterile rubber surgical gloves					
ISO 10282 Technical (Corri 2005-06	N	Specification; Technical Corrigendum_1					
			Ophthalmic optics Semi-finished spectacle lens					
			blanks Part_1: Specifications for single-vision					
ISO 10322-1	2006-02	N	and multifocal lens blanks					
			Ophthalmic optics Semi-finished spectacle lens					
			blanks Part_2: Specifications for progressive					
ISO 10322-2	2006-02	N	power lens blanks					

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100 40222	1001 11	N.	Dental rotary instruments; bore diameters for discs and wheels					
ISO 10323	1991-11	N	Prosthetics Structural testing of lower-limb prostheses					
ISO 10328	2006-10	N	Requirements and test methods					
			Implants for surgery Malleable wires for use as					
ISO 10334	1994-08	N	sutures and other surgical applications					
ISO 10341	2009-07	N	Ophthalmic instruments Refractor heads					
ISO 10342	2010-06	N	Ophthalmic instruments Eye refractometers	Υ	N	JIS_T_7319:2011	29.07.2011	610
ISO 10343	2009-07	N	Ophthalmic instruments Ophthalmometers	Р	N	JIS_T_7318:2002	01.08.2002	610
			Dentistry Contents of technical file for dental implant					
ISO 10451	2010-06	N	systems	Υ	N			AP36
						JIS T 6517:2011	2011/7/29	232
ISO 10477	2004-10	N	Dentistry Polymer-based crown and bridge materials	Р	N	JIS T 6518:2011	2011/7/29	231
			Pressure regulators for use with medical gases					
			Part_1: Pressure regulators and pressure					
ISO 10524-1	2006-02	N	regulators with flow-metering devices					
100 40504.0	0005.05		Pressure regulators for use with medical gases					
ISO 10524-2	2005-05	N	Part_2: Manifold and line pressure regulators					
			Pressure regulators for use with medical gases					
ISO 10524-3	2005-05	N	Part_3: Pressure regulators integrated with cylinder valves					
150 10524-3	2005-05	IN	Pressure regulators for use with medical gases -		 			
ISO 10524-4	2008-06	N	Part_4: Low-pressure regulators					
130 10324-4	2000-00	IN	Hoists for the transfer of disabled persons					
ISO 10535	2006-12	N	Requirements and test methods					
130 10333	2000-12	IN	Technical systems and aids for disabled or handicapped					
			persons Wheelchair tiedown and occupant-restraint					
			systems Part_1: Requirements and test methods for all					
ISO 10542-1	2001-07	N	systems					
			T					
			Technical systems and aids for disabled or handicapped persons Wheelchair tiedown and occupant-restraint					
ISO 10542-2	2001-07	N	systems Part_2: Four-point strap-type tiedown systems					
130 10342-2	2001-07	IN	Systems anz. r our point on up type abdomin systems					
			Technical systems and aids for disabled or handicapped					
			persons Wheelchair tiedown and occupant-restraint					
ISO 10542-3	2005-02	N	systems Part_3: Docking-type tiedown systems					
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			Technical systems and aids for disabled or handicapped persons - Wheelchair tiedown and occupant-restraint					
ISO 10542-4	2004-09	N	systems - Part 4: Clamp-type tiedown systems					
150 10542-4	2004-09	IN	systems1 art_4. Olamp-type fledown systems					
			Technical systems and aids for disabled or handicapped					
			persons Wheelchair tiedown and occupant-restraint					
ISO 10542-5	2004-04	N	systems Part_5: Systems for specific wheelchairs					
			Sterile, single-use intravascular catheters Part_1:					
ISO 10555-1	1995-06	N	General requirements	Υ	N			AP3,AP23,AP39,AP40,AP43
			Sterile, single-use intravascular catheters Part_1:					
ISO 10555-1 AMD 1	1999-07	N	General requirements; Amendment_1	Р	N	JIS T 3268:2012	01.03.2012	823
			Sterile, single-use intravascular catheters Part_1:					
ISO 10555-1 AMD 2	2004-05	N	General requirements; Amendment_2	Р	N	JIS T 3268:2012	01.03.2012	823
			Sterile, single-use intravascular catheters Part_2:					
ISO 10555-2	1996-06	N	Angiographic catheters	Р	N	JIS T 3268:2012	01.03.2012	823

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		Sterile, single-use intravascular catheters Part_2:					
ISO 10555-2 Technical Co 2002-06	N	Angiographic catheters; Technical Corrigendum_1					
ISO 10555-3 1996-06	N	Sterile, single-use intravascular catheters Part_3: Central venous catheters					
150 10555-3	IN	Sterile, single-use intravascular catheters Part_3:					
		Central venous catheters; Technical					
ISO 10555-3 Technical Co 2002-06	N	Corrigendum 1					
130 10333-3 Technical Co 2002-00	IN	Sterile, single-use intravascular catheters - Part 4:					
ISO 10555-4 1996-06	N	Balloon dilatation catheters	Υ	N			AP3•AP23
1990-00	IN	Sterile, single-use intravascular catheters - Part 4:		IN			AI 3 AI 20
		Balloon dilatation catheters; Technical					
ISO 10555-4 Technical Co 2002-06	N	Corrigendum 1					
100 1000 1 100 mman 00 2002 00		Sterile, single-use intravascular catheters Part_5:			JIS T 3249:2011	2011/7/29	93
ISO 10555-5 1996-06	N	Over-needle peripheral catheters	Р	N	JIS T 3223:2011	2011/7/29	73 · 631 · 634
1990-00	IN	Over riceale periprieral satisficies	•	IN IN	010 1 0220.2011	2011/1/20	73 031 034
		Sterile, single-use intravascular catheters Part_5:			JIS T 3249:2011	2011/7/29	93
ISO 10555-5 AMD 1 1999-01	N	Over-needle peripheral catheters; Amendment_1	Р	N	JIS T 3249.2011 JIS T 3223:2011	2011/7/29	73 · 631 · 634
100 10000-3 AIVID 1 1999-01	IN .	Sterile, single-use intravascular catheters Part_5:	Г	IN	0.0 . 0220.20	2011/1/20	73 031 034
		Over-needle peripheral catheters; Technical					
ISO 10555-5 Technical Co 2002-06	N	Corrigendum 1					
130 10333-3 Technical Co 2002-00	IN	Dental equipment High- and medium-volume suction					
ISO 10637 1999-08	N	systems	Р	N	JIS T 5801:2005	25.03.2005	406
100 1001		Dentistry Powered polymerization activators Part_1:	· ·	.,	0.0 1 0001.2000	20.00.2000	1.00
ISO 10650-1 2004-11	N	Quartz tungsten halogen lamps					
		Dentistry Powered polymerization activators Part_2:					
ISO 10650-2 2007-09	N	Light-emitting diode (LED) lamps					
		Lung ventilators for medical use Particular					
		requirements for basic safety and essential					
		performance Part_2: Home care ventilators for					
ISO 10651-2 2004-07	N	ventilator-dependent patients					
		Lung ventilators for medical use Part_3:					
		Particular requirements for emergency and					
ISO 10651-3 1997-01	N	transport ventilators					
100 10051 1		Lung ventilators Part_4: Particular requirements					
ISO 10651-4 2002-03	N	for operator-powered resuscitators					
		Lung ventilators for medical use Particular requirements for basic safety and essential					
		performance Part_5: Gas-powered emergency					
ISO 10651-5 2006-02	N	resuscitators					
2000-02	IN	Lung ventilators for medical use Particular					
		requirements for basic safety and essential					
		performance Part_6: Home-care ventilatory					
ISO 10651-6 2004-07	N	support devices					
		Ophthalmic optics Spectacle frames and					
	1	sunglasses electronic catalogue and identification					
	1	Part_1: Product identification and electronic					
ISO 10685-1 2011-12	N	catalogue product hierarchy					
ISO 10873 2010-09	N	Dentistry Denture adhesives					
		Optics and optical instruments Operation					
		microscopes Part_1: Requirements and test					
ISO 10936-1 2000-06	N	methods					

			Optics and photonics Operation microscopes					
			Part 2: Light hazard from operation microscopes					
ISO 10936-2	2010-01	N	used in ocular surgery					
ISO 10938	1998-05	N	Ophthalmic instruments Chart projectors					
ISO 10939	2007-02	N	Ophthalmic instruments Slit-lamp microscopes					
ISO 10940	2009-08	N	Ophthalmic instruments Fundus cameras					
100 100 10								
ISO 10942	2006-06	N	Ophthalmic instruments Direct ophthalmoscopes					
ISO 10943	2011-08	N	Ophthalmic instruments Indirect ophthalmoscopes					
ISO 10943	2009-08	N N	Ophthalmic instruments - Synoptophores					
100 10344	2009-00	IN .	Caps made of aluminium-plastics combinations for					
			infusion bottles and injection vials Requirements					
ISO 10985	2009-02	N	and test methods					
			Biological evaluation of medical devices Part_1:					
			Evaluation and testing within a risk management					
ISO 10993-1	2009-10	N	process	Р	Р	JIS T 0993-1:2012	01.03.2012	See # 2 (429) on Sheet1
			Biological evaluation of medical devices Part_1:					
			Evaluation and testing within a risk management					
ISO 10993-1 Technic	al Co 2010-06	N	process; Technical Corrigendum_1	Р	Р	JIS T 0993-1:2012	01.03.2012	See # 2 (429) on Sheet1
			Biological evaluation of medical devices Part_10:					
ISO 10993-10	2010-08	N	Tests for irritation and skin sensitization Biological evaluation of medical devices Part_11:	Y	N			MHLW Ministerial Notice
100 40000 44	0000 00	N	Tests for systemic toxicity	Υ				MHLW Ministerial Notice
ISO 10993-11	2006-08	N	resis for systemic toxicity	Y	N			MINE VV MINISTERIAL NOTICE
			Biological evaluation of medical devices Part_12:					
ISO 10993-12	2007-11	N	Sample preparation and reference materials	Υ	N			MHLW Ministerial Notice
100 10333 12	2007 11			<u> </u>	111			IVII IEVV IVIII II OLOGO
			Biological evaluation of medical devices Part_13:					
			Identification and quantification of degradation					
ISO 10993-13	2010-06	N	products from polymeric medical devices					
			Biological evaluation of medical devices Part_14:					
			Identification and quantification of degradation					
ISO 10993-14	2001-11	N	products from ceramics	Υ	N			GL4,5,6,7,8
			Biological evaluation of medical devices Part_15:					
			Identification and quantification of degradation					
ISO 10993-15	2000-12	N	products from metals and alloys	Υ	N			GL4,5,6,7,8
			Biological evaluation of medical devices Part_16:					
100 10000 10	0040.00		Toxicokinetic study design for degradation products					
ISO 10993-16	2010-02	N	and leachables Biological evaluation of medical devices Part_17:					
			Establishment of allowable limits for leachable					
ISO 10993-17	2002-12	N	substances	Υ	N			GL4.5.6.7.8
10000 17	2002 12	14	Biological evaluation of medical devices Part_18:		14			52.,5,5,1,5
ISO 10993-18	2005-07	N	Chemical characterization of materials					
	2000 01		Biological evaluation of medical devices Part_2:		1			
ISO 10993-2	2006-07	N	Animal welfare requirements	Υ	N			MHLW Ministerial Notice

			Biological evaluation of medical devices Part_3:					
ISO 10993-3	2003-10	N	Tests for genotoxicity, carcinogenicity and reproductive toxicity	Υ	N			MHLW Ministerial Notice
	2000 10			· ·	.,			im iz v immeteriar v teaec
ISO 10993-4	2002-10	N	Biological evaluation of medical devices Part_4: Selection of test for interactions with blood	Υ	N			MHLW Ministerial Notice
100 40000 4 440 4	0000 07		Biological evaluation of medical devices Part_4: Selection of tests for interactions with blood					NALIL VAZ BAGOGIA A OSI ALI NILAGIA A
ISO 10993-4 AMD 1	2006-07	N	Biological evaluation of medical devices - Part 5:	Y	N			MHLW Ministerial Notice
ISO 10993-5	2009-06	N	Tests for in vitro cytotoxicity	Υ	N			MHLW Ministerial Notice
100 10995-5	2009-00	11	Biological evaluation of medical devices Part_6:		11			WITTERV WITTISTETIAL TVOLICE
ISO 10993-6	2007-04	N	Tests for local effects after implantation	Υ	N			MHLW Ministerial Notice
			Biological evaluation of medical devices Part_7:					
ISO 10993-7	2008-10	N	Ethylene oxide sterilization residuals	Υ	N	JIS T 0993-7:2012	01.03.2012	MHLW Ministerial Notice
			Biological evaluation of medical devices Part_7:					
			Ethylene oxide sterilization residuals; Technical					
ISO 10993-7 Technica	Il Co 2009-11	N	Corrigendum_1	Υ	N	JIS T 0993-7:2012	01.03.2012	MHLW Ministerial Notice
			Biological evaluation of medical devices Part_9: Framework for identification and quantification of					
ISO 10993-9	2009-12	N	potential degradation products	Υ	N			MHLW Ministerial Notice
130 10993-9	2009-12	IN	Prefilled syringes; part_1: glass cylinders for dental	ı	IN IN			IVII IEVV IVIII II Steriai Notice
ISO 11040-1	1992-11	N	local anaesthetic cartridges					
100 110 10 1	1002 11		Prefilled syringes Part_2: Plunger stoppers for					
ISO 11040-2	2011-04	N	dental local anaesthetic cartridges Prefilled syringes Part_3: Seals for dental local					
			Prefilled syringes Part_3: Seals for dental local anaesthetic cartridges					
ISO 11040-3	2012-01	N	5					
ISO 11040-4	2007-02	N	Prefilled syringes Part_4: Glass barrels for injectables					
130 11040-4	2007-02	IN	Prefilled syringes - Part 5: Plunger stoppers for					
ISO 11040-5	2012-01	N	injectables					
						UO T 0007-0040	0040/0/4	761 · AP47
						JIS T 3267:2013 JIS T 3262:2012	2013/3/1 2012/10/1	630· 391
						JIS T 3261:2012	2012/10/1	388 656 665
				Р		JIS T 3260:2012	2012/10/1	665· 396
ISO 11070	1998-05	N	Sterile single-use intravascular catheter introducers	Υ	N			AP41· AP42
			Health informatics Point-of-care medical device					
ISO 11073-90101	2008-01	N	communication Part_90101: Analytical instruments Point-of-care test					
150 11073-90101	2008-01	IN IN	r onit-or-care test					
			Health informatics Standard communication protocol					
ISO 11073-91064	2009-05	N	Part_91064: Computer-assisted electrocardiography					
			Sterilization of health care products Ethylene					
			oxide Part_1: Requirements for development,					MHLW Ministerial Notice
ISO 11125 1	2007.05	N.	validation and routine control of a sterilization process for medical devices	V	N	IIC T 0004 4,2040	25.02.2040	
ISO 11135-1	2007-05	N	Sterilization of health care products Radiation	Υ	N	JIS T 0801-1:2010	25.02.2010	AP41,42
		1	Part_1: Requirements for development, validation					
			and routine control of a sterilization process for					MHLW Ministerial Notice
ISO 11137-1	2006-04	N	medical devices	Υ	N	JIS T 0806-1:2010	25.02.2010	AP41,42

	1		Otavilla stick of health and another Dedication					MHLW Ministerial Notice
			Sterilization of health care products Radiation	.,				
ISO 11137-2	2012-03	N	Part_2: Establishing the sterilization dose	Y	N	JIS T 0806-2:2010	25.02.2010	AP41,42
			Sterilization of health care products Radiation					
ISO 11137-3	2006-04	N	Part_3: Guidance on dosimetric aspects	Y	N			AP41 • AP42
			Sterilization of health care products Biological					
ISO 11138-1	2006-07	N	indicators Part_1: General requirements					
			Sterilization of health care products Biological					
			indicators Part_2: Biological indicators for					
ISO 11138-2	2006-07	N	ethylene oxide sterilization processes					
			Sterilization of health care products Biological					
			indicators Part_3: Biological indicators for moist					
ISO 11138-3	2006-07	N	heat sterilization processes					
			Sterilization of health care products Biological					
			indicators Part_4: Biological indicators for dry					
ISO 11138-4	2006-07	N	heat sterilization 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					
ISO 11140-1	2005-07	N	indicators Part_1: General requirements					
			Sterilization of health care products Chemical					
			indicators Part_3: Class_2 indicator systems for					
100 11110 0	0007.00		use in the Bowie and Dick-type steam penetration					
ISO 11140-3	2007-03	N	test					
			Sterilization of health care products Chemical					
			indicators Part_3: Class 2 indicator systems for					
100 111 10 0 T 1 1 0	0007.44		use in the Bowie and Dick-type steam penetration					
ISO 11140-3 Technical Co	2007-11	N	test; Technical Corrigendum_1					
			Sterilization of health care products Chemical					
			indicators Part_4: Class_2 indicators as an					
100 44440 4	0007.00	N.	alternative to the Bowie and Dick-type test for					
ISO 11140-4	2007-03	N	detection of steam penetration					
			Sterilization of health care products Chemical					
ICO 44440 F	2007.02	N.	indicators Part_5: Class_2 indicators for Bowie and Dick-type air removal tests					
ISO 11140-5	2007-03	N	Dentistry Amalgam separators					
ISO 11143	2008-07	N	Dental equipment Connections for supply and waste					
ISO 11144	1995-05	N	lines					
150 11144	1995-05	IN	inios					
ISO 11156	2011-07	N	Packaging Accessible design General requirements					
100 11100	2011-07	18	Single-use medical examination gloves - Part 1:					
			Specification for gloves made from rubber latex or					
ISO 11193-1	2008-09	N	rubber solution					
100 11130-1	2000-09	IN	Single-use medical examination gloves Part_2:					
			Specification for gloves made from poly(vinyl					
ISO 11193-2	2006-11	N	chloride)					
100 11180-2	2000-11	IN	Gas mixers for medical use Stand-alone gas					
ISO 11195	1995-10	N	mixers					
ISO 11195 ISO 11197	2004-12	N N	Medical supply units		1			
30 11181	2004-12	IN	iviculcal supply utilis		1			

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Pen-injectors for medical use, - Part 3, Finished contributes and test methods Pen-injectors for medical use, - Part 4,							 	
Peni-injectors for medical use Part_4				Pen-injectors for medical use Part_3: Finished				
So 11608-4 2006-03	ISO 11608-3	2000-12	N	cartridges; Requirements and test methods				
				Pen-injectors for medical use Part_4:				
Societion Soci								
ISO 1169 2010-99 N marking	ISO 11608-4	2006-03	N					
SO 11663 2009-04 N	100 44000	0040.00						
ISO 11683 2009-04	ISO 11609	2010-09	N	9				
So 11683 1997-10 N Paskaging - Tactile warnings of danger - Requirements	100 44000	0000 04	N					
ISO 11712 2009-05 N Surplangeal almoys and connectors Sterifization of medical devices — Microbiological methods — Part 1: Determination of a population of microorganisms on products Sterifization of medical devices — Microbiological methods — Part 1: Determination of a population of microorganisms on products Sterifization of medical devices — Microbiological methods — Part 1: Determination of a population of microorganisms on products. Technical Co. 2007-05 N Corrigendum_1 ISO 11737-1 Technical Co. 2007-05 N Corrigendum_1 ISO 11737-2 2009-11 N Sterifization of medical devices — Microbiological methods — Part 2: Tests of sterifity performed in the definition, validation and maintenance of a sterifization process Lasers and laser-related equipment _ Test method and classification for the laser resistance of surgical drapes and/or patient protective covers _ Part 1: Primary ignition and penetration Lasers and laser-related equipment _ Test method and classification for the laser resistance of surgical drapes and/or patient protective covers _ Part 1: Primary ignition and penetration Lasers and laser-related equipment _ Test method and classification for the laser resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser-resistance of surgical drapes and/or patient-protective covers _ Part 1: Test method and classification for the laser resistance of surgical drapes a	150 11663	2009-04	IN .	related theraples				
ISO 11712 2009-05 N Supratargoeal airways and connectors Sterification of medical devices — Microbiological methods — Part .1: Determination of a population of microorganisms on products Sterification of medical devices — Microbiological methods — Part .1: Determination of a population of microorganisms on products — Technical Co. 2007-05 N Congendum .1 ISO 11737-1 Technical Co. 2007-05 N Congendum .1 ISO 11737-2 2009-11 N Sterification of medical devices — Microbiological methods — Part .2: Tests of sterifity performed in the definition, validation and maintenance of a sterification process Lasers and laser-related equipment — Test method and classification for the laser resistance of surgical drapes and/or patient protective covers — Part .1: Permany ignition and penetration Lasers and laser-related equipment — Test method and classification for the laser resistance of surgical drapes and/or patient protective covers — Part .1: Permany ignition and penetration Lasers and laser-related equipment — Test method and classification for the laser resistance of surgical drapes and/or patient protective covers — Part .1: Permany ignition and penetration Sol 11810-2 2007-05 N Primary ignition and penetration Sol 11810-1 2002-10 N Part 2. Secondary ignition of microbion in a mail and (MIRE technique) Sol 11804-1 2002-10 N International Part .1: Whole-product testing Definition of the structure of the part .1: Technique using a microphone in a real set of IRIE technique) Sol 11804-1 1996-11 N University — Intraocolural lenses and contact lenses are products — Information supplied by the manufacturer Ophthalmic optics — Contact lenses — Part .1: Whole-product testing Ophthalmic optics — Contact lenses — Part .1: Whole-product lenses — Ophthalmic implants — Intraocolural lenses — Optical penese — Ophthalmic implants — Intraocolural lenses — Ophthalmic implants — Intraocolural lenses — Ophthalmic implants — Intraocolural lenses — Part .1: Whole-product lenses — Ophthalmic implants — Intraocolural lenses — Par	ISO 11683	1997-10	N	Packaging Tactile warnings of danger Requirements				
ISO 11712 2009-05 N Supralaryngeal airways and connectors		1001.10		Anaesthetic and respiratory equipment -				
Sterilization of medical devices _ Microbiological methods _ Part_ 1: Determination of a population of microorganisms on products	ISO 11712	2009-05	N					
ISO 11737-1					-			
Sterilization of medical devices Microbiological methods_ Part_1: Determination of a population of microorganisms on products; Technical Co 2007-05 N Corrigendum_1								
methods_Part_1: Determination of a population of microorganisms on products; Technical configuration of microorganisms on products; Technical configuration of medical devices_Microbiological methods_Part_2: Tests of stenity performed in the definition, validation and maintenance of a stenity performed in the definition, validation and maintenance of a stenity performed and classification for the laser resistance of surgical drapes and/or patient profective covers_Part_1: SO 11810-1	ISO 11737-1	2006-04	N	microorganisms on products				
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Sterilization of medical devices _ Microbiological methods_Part_2: Tests of sterility performed in the definition, validation and maintenance of a sterilization process				microorganisms on products; Technical				
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Ophthalmic implants Intraocular lenses	ISO 11979-1	2006-07	N					
	100 11070-1	2000-01	111				1	+
	ISO 11979-10	2006-08	N					
Ophthalmic implants - Intraocular lenses - Part_2:		2000 00						
ISO 11979-2 1999-12 N Optical properties and test methods Y N	ISO 11979-2	1999-12	N		Υ	N		AP2
Ophthalmic implants Intraocular lenses Part_2:								
Optical properties and test methods; Technical						1		
ISO 11979-2 Technical Co 2003-11 N Corrigendum_1	ISO 11979-2 Technical	l Co 2003-11	N	Corrigendum_1				

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			Ophthalmic implants Intraocular lenses Part_3:				
ISO 11979-3	2006-05	N	Mechanical properties and test methods	Y	N		AP2
			Ophthalmic implants Intraocular lenses Part_4:				
ISO 11979-4	2008-12	N	Labelling and information	Υ	N		AP2
			Ophthalmic implants Intraocular lenses Part_5:				
ISO 11979-5	2006-06	N	Biocompatibility	Υ	N		AP2
			Ophthalmic implants Intraocular lenses Part_6:				
ISO 11979-6	2007-07	N	Shelf-life and transport stability	Υ	N		AP2
			Ophthalmic implants Intraocular lenses Part_7:				
ISO 11979-7	2006-05	N	Clinical investigations				
			Ophthalmic implants Intraocular lenses Part_7:				
ISO 11979-7 AMD 1	2012-01	N	Clinical investigations; Amendment 1				
	20.2 0.		Ophthalmic implants - Intraocular lenses - Part 8:				
ISO 11979-8	2006-07	N	Fundamental requirements				
100 11070 0	2000 07		T disamental requirements				
			Ophthalmic implants Intraocular lenses Part_8:				
ISO 11979-8 AMD 1	2011-05	N	Fundamental requirements; Amendment_1				
100 11070 074012 1	2011 00	1	Ophthalmic implants - Intraocular lenses - Part 9:				
ISO 11979-9	2006-09	N	Multifocal intraocular lenses				
100 11373 3	2000 03	11	Ophthalmic optics - Contact lenses and contact				
			lens care products Guidance for clinical				
ISO 11980	2009-10	N	investigations				
150 11980	2003-10	IN	Ophthalmic optics - Contact lenses and contact				
			lens care products Determination of physical				
			compatibility of contact lens care products with				
ISO 11981	2009-07	N	contact lenses				
130 11981	2009-07	IN	Ophthalmic optics - Contact lenses - Ageing by				
			exposure to UV and visible radiation (in vitro				
ISO 11985	1997-12	N	method)				
130 11985	1997-12	IN	Ophthalmic optics - Contact lenses and contact				
			lens care products Determination of preservative				
ISO 11986	2010-11	N	uptake and release				
130 11966	2010-11	IN	Ophthalmic optics Contact lenses				
ISO 11987	1997-12	N	Determination of shelf-life	Υ	N		AP1
130 11967	1997-12	IN	Ophthalmic optics Contact lenses	T	IN		API
			Determination of shelf-life; Technical				
ISO 11987 Technical Corr	1000.04	N	Corrigendum 1				
130 11967 Technical Con	1990-04	IN	Lasers and laser-related equipment -				
			Determination of laser resistance of tracheal tubes				
ISO 11990-1	2011-08	N	Part 1: Tracheal tube shaft				
150 11990-1	2011-08	IN	Lasers and laser-related equipment -				
			Determination of laser resistance of tracheal tubes				
100 44000 0	0040.07	N.	_				
ISO 11990-2	2010-07	N	Part_2: Tracheal tube cuffs Health informatics Digital imaging and communication in				
			medicine (DICOM) including workflow and data				
ISO 12052	2006-11	N	management				
		1.	Acoustics Procedures for the measurement of real-ear			†	
ISO 12124	2001-03	N	acoustical characteristics of hearing aids				
			Implants for surgery - Mechanical testing of				
	1		implantable spinal devices Fatigue test method				
			for spinal implant assemblies using an anterior				
ISO 12189	2008-05	N	support				
	1-200 00		1			1	

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			Medical gloves made from natural rubber latex				
			Determination of water-extractable protein using				
ISO 12243	2003-10	N	the modified Lowry method				
			Tissue paper and tissue products Part_1: General				
ISO 12625-1	2011-08	N	guidance on terms				
			Tissue paper and tissue products Part_12:				
			Determination of tensile strength of perforated lines				
ISO 12625-12	2010-01	N	Calculation of perforation efficiency				
			Tissue paper and tissue products - Part 3: Determination				
100 40005 0	0005.04		of thickness, bulking thickness and apparent bulk density				
ISO 12625-3	2005-04	N	Tissue paper and tissue products - Part 4: Determination				
			of tensile strength, stretch at break and tensile energy				
ISO 12625-4	2005-04	N	absorption				
100 12025 4	2003-04	14	Tissue paper and tissue products Part_5: Determination		+		
ISO 12625-5	2005-04	N	of wet tensile strength				
100 12020 0	2000 04	- ''	Tissue paper and tissue products Part_6: Determination				
ISO 12625-6	2005-02	N	of grammage				
100 12020 0	2000 02	.,	Tissue paper and tissue products Part_7: Determination				
ISO 12625-7	2007-03	N	of optical properties				
			Tissue paper and tissue products Part_8: Water-				
			absorption time and water-absorption capacity, basket-				
ISO 12625-8	2010-12	N	immersion test method				
			Tissue paper and tissue products Part_9: Determination				
ISO 12625-9	2005-05	N	of ball burst strength				
			Ophthalmic optics Contact lenses				
ISO 12864	1997-12	N	Determination of scattered light				
ISO 12865	2006-07	N	Ophthalmic instruments Retinoscopes				
ISO 12866	1999-06	N	Ophthalmic instruments Perimeters				
			Ophthalmic instruments - Perimeters;				
ISO 12866 AMD 1	2008-11	N	Amendment 1				
ISO 12867	2010-06	N	Ophthalmic instruments Trial frames				
			Ophthalmic optics Spectacle frames				
ISO 12870	2004-08	N	Requirements and test methods				
			Implants for surgery Retrieval and analysis of				
ISO 12891-1	2011-05	N	surgical implants Part_1: Retrieval and handling				
			Retrieval and analysis of surgical implants				
			Part 2: Analysis of retrieved metallic surgical				
ISO 12891-2	2000-02	N	implants				
			Retrieval and analysis of surgical implants -				
			Part 3: Analysis of retrieved polymeric surgical				
ISO 12891-3	2000-02	N	implants				
100 12001 0	2000 02		Retrieval and analysis of surgical implants				
			Part_4: Analysis of retrieved ceramic surgical				
ISO 12891-4	2000-02	N	implants				
	2000 02	 	Health informatics Service architecture Part_1:		<u> </u>	1	
ISO 12967-1	2009-08	N	Enterprise viewpoint				
		İ	Health informatics Service architecture Part_2:				
ISO 12967-2	2009-08	N	Information viewpoint				
			Health informatics Service architecture Part_3:				
ISO 12967-3	2009-08	N	Computational viewpoint				
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			Ophthalmic optics - Contact lens care products -					
ISO 13212	2011-05	N	Guidelines for determination of shelf-life		1			
ISO 13294	1997-05	N	Dental handpieces Dental air-motors	P	N	JIS T 5908:2012	01.03.2012	151 · 486
ISO 13295	2007-07	N	Dentistry - Mandrels for rotary instruments					
100 10200	2001-01	11	Implants for surgery Ceramic materials based on					
ISO 13356	2008-06	N		Υ	l N			AP36
100 10000	2000 00	.,	yttria-stabilized tetragonal zirconia (Y-TZP) Periodontal curettes, dental scalers and excavators	•	1			711 00
ISO 13397-1	1995-12	N	Part_1: General requirements					
			Dentistry Periodontal curettes, dental scalers and					
ISO 13397-2	2005-06	N	excavators Part_2: Periodontal curettes of Gr-type					
100 10007 0	4000 00		Periodontal curettes, dental scalers and excavators Part 3: Dental scalers - H-type					
ISO 13397-3	1996-09	N	Periodontal curettes, dental scalers and excavators					
ISO 13397-4	1997-12	N	Part_4: Dental excavators Discoid type					
100 10001 4	1007 12	11	Surgical and dental hand instruments Determination of					
			resistance against autoclaving, corrosion and thermal					
ISO 13402	1995-08	N	exposure					
			Prosthetics and orthotics Categorization and description					
ISO 13404	2007-07	N	of external orthoses and orthotic components Prosthetics and orthostics - Classification and description					
			of prosthetic components Part_1: Classification of					
ISO 13405-1	1996-10	N	prosthetic components					
100 10 100 1	1000 10		Prosthetics and orthostics Classification and description					
			of prosthetic components Part_2: Description of lower-					
ISO 13405-2	1996-10	N	limb prosthetic components					
			Prosthetics and orthostics Classification and description					
100 40 40 5	1000 10		of prosthetic components Part_3: Description of upper- limb prosthetic components					
ISO 13405-3	1996-10	N	· · · · · · · · · · · · · · · · · · ·					
100 40 400 4	0000 00	N.	Aseptic processing of health care products					
ISO 13408-1	2008-06	N	Part_1: General requirements Aseptic processing of health care products					
ISO 13408-2	2003-03	N	Part 2: Filtration					
130 13406-2	2003-03	IN	Aseptic processing of health care products -					
ISO 13408-3	2006-09	N	Part_3: Lyophilization					
100 10400-0	2000-03	IN	Aseptic processing of health care products					
ISO 13408-4	2005-11	N	Part 4: Clean-in-place technologies					
100 10100 1	2000	.,	Aseptic processing of health care products -					
ISO 13408-5	2006-11	N	Part_5: Sterilization in place					
			Aseptic processing of health care products					
ISO 13408-6	2005-06	N	Part_6: Isolator systems					
			Medical devices Quality management systems					MHLW Ministerial Ordinance No.
ISO 13485	2003-07	N	Requirements for regulatory purposes	Y	N	JIS Q 13485:2005	01.10.2005	169,2004
			Medical devices Quality management systems					
	1		Requirements for regulatory purposes; Technical					
ISO 13485 Technical Cor	ri 2009-08	N	Corrigendum_1					
100 40000 4	0000 00	l N	Health informatics Electronic health record communication - Part 1: Reference model					
ISO 13606-1	2008-02	N	Health informatics Electronic health record			+		
			communication Part_2: Archetype interchange					
ISO 13606-2	2008-12	N	specification		1			

			Health informatics Electronic health record		1		1	T
			communication - Part 3: Reference archetypes and term					
ISO 13606-3	2009-02	N	lists					
100 10000 0	2000 02	- ''	Health informatics - Electronic health record					
ISO 13606-5	2010-03	N	communication Part_5: Interface specification					
ISO 13666	1998-08	N	Ophthalmic optics Spectacle lenses Vocabulary Dentistry Reversible-irreversible hydrocolloid					
ISO 13716	1999-05	N	impression material systems					
			Implants for surgery Hydroxyapatite Part_1:					
ISO 13779-1	2008-10	N	Ceramic hydroxyapatite					
			Implants for surgery Hydroxyapatite Part_2:					
ISO 13779-2	2008-10	N	Coatings of hydroxyapatite					
			Implants for surgery Hydroxyapatite Part_3:					
100 40770 0	0000 00	N.	Chemical analysis and characterization of					
ISO 13779-3	2008-02	N	crystallinity and phase purity					
100 40770 4	2002.05	N	Implants for surgery Hydroxyapatite Part_4: Determination of coating adhesion strength					
ISO 13779-4	2002-05	IN	Poly(L-lactide) resins and fabricated forms for			+		
ISO 13781	1997-02	N	surgical implants In vitro degradation testing					
130 13761	1997-02	IN	surgical implants in vitro degradation testing			+		
			Implants for surgery - Metallic materials -					
ISO 13782	1996-12	N	Unalloyed tantalum for surgical implant applications					
ISO 13897	2003-02	N	Dentistry - Amalgam capsules					
100 10007	2000 02	- ''	3,					
ISO 13897 Technical C	orriger 2003-12	N	Dentistry Amalgam capsules; Technical Corrigendum_1					
			Pen systems - Part 1: Glass cylinders for pen-					
ISO 13926-1	2004-11	N	injectors for medical use					
			Pen systems Part_2: Plunger stoppers for pen-					
ISO 13926-2	2011-04	N	injectors for medical use					
			Concentrates for haemodialysis and related					
ISO 13958	2009-04	N	therapies					
ISO 13959	2009-04	N	Water for haemodialysis and related therapies					
			Cardiovascular implants and extracorporeal					
ISO 13960	2010-07	N	systems Plasmafilters					
			Clinical investigation of medical devices for human					MHLW Ministerial Ordinance No.
ISO 14155	2011-02	N	subjects Good clinical practice	Υ	N			36,2005
			Clinical investigation of medical devices for human					
			subjects Good clinical practice; Technical					
ISO 14155 Technical	l Corri 2011-07	N	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,					
100 44400	0044.07	N.	development, validation and routine control of a					
ISO 14160	2011-07	N	sterilization process for medical devices					
			Sterilization of health care products Biological					
ISO 14161	2009-09	N	indicators Guidance for the selection, use and interpretation of results					
ISO 14161	2009-09	N N	Dentistry Polymer-based die materials					
130 14233	2003-03	IN	Domining i digitici-basca die materials					

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			Implants for surgery Wear of total hip-joint				
			prostheses Part_1: Loading and displacement				
			parameters for wear-testing machines and				
ISO 14242-1	2012-01	N	corresponding environmental conditions for test	Υ	N		GL1
			Implants for surgery Wear of total hip joint				
ISO 14242-2	2000-09	N	prostheses Part_2: Methods of measurement	Y	N		GL1
			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				
ISO 14242-3	2009-03	N	conditions 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				
ISO 14243-1	2009-11	N	conditions for test	Υ	N		GL2
			Implants for surgery Wear of total knee-joint				
ISO 14243-2	2009-11	N	prostheses Part_2: Methods of measurement	Y	N		GL2
			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	Y	N		GL2
			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 Techni		N	environmental conditions for test				
ISO 14356	2003-03	N	Dentistry Duplicating material				
			Tracheal tubes designed for laser surgery				
			Requirements for marking and accompanying				
ISO 14408	2005-06	N	information				
			Ophthalmic optics Contact lenses and contact				
ISO 14534	2011-04	N	lens care products Fundamental requirements				
			Non-active surgical implants Implants for				
ISO 14602	2010-04	N	osteosynthesis Particular requirements				
			Non-active surgical implants Mammary implants				
ISO 14607	2007-02	N	Particular requirements				
			Non-active surgical implants General				
ISO 14630	2008-01	N	requirements	Y	N		AP27 • 28 • AP42
			Implants for surgery Active implantable medical				
			devices Part_1: General requirements for safety,				
			marking and for information to be provided by the	.,			
ISO 14708-1	2000-11	N	manufacturer	Y	N		AP18
			Implants for surgery Active implantable medical	.,			
ISO 14708-2	2005-10	N	devices Part_2: Cardiac pacemakers	Y	N		AP18
			landanta fan armana. Astira insalantahi 1911				
100 44700 0	2000 44	N.1	Implants for surgery Active implantable medical				
ISO 14708-3	2008-11	N	devices Part_3: Implantable neurostimulators				
			Implants for surgery Active implantable medical				
ISO 14708-4	2008-11	N	devices Part_4: Implantable infusion pumps				
130 14/00-4	2000-11	IN	uevices rait_4. impiantable iniusion pumps				

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			Implants for surgery Active implantable medical					
ISO 14708-5	2010-02	N	devices Part_5: Circulatory support devices					
			Implants for surgery Active implantable medical					
			devices Part_6: Particular requirements for active					
			implantable medical devices intended to treat					
			tachyarrhythmia (including implantable					
ISO 14708-6	2010-03	N	defibrillators)					
			Ophthalmic optics Contact lens care products					
			Microbiological requirements and test methods for					
ISO 14729	2001-04	N	products and regimens for hygienic management of contact lenses					
130 14729	2001-04	IN	Ophthalmic optics Contact lens care products					
			Microbiological requirements and test methods for					
			products and regimens for hygienic management of					
ISO 14729 AMD 1	2010-10	N	contact lenses; Amendment 1					
	2010 10		oonidat ionidaa, runandina					
			Ophthalmic optics Contact lens care products					
			Antimicrobial preservative efficacy testing and					
ISO 14730	2000-09	N	guidance on determining discard date					
			Dentistry Implants Dynamic fatigue test for					
ISO 14801	2007-11	N	endosseous dental implants	Y	N			AP36
			Implants for surgery Total knee-joint prostheses					
100 44070 4	0000 00		Part_1: Determination of endurance properties of					
ISO 14879-1	2000-06	N	knee tibial trays	Y	N			GL2
			Ophthalmic optics Spectacle lenses Fundamental requirements for uncut finished					
ISO 14889	2003-05	N	lenses					
130 14009	2003-03	IN	lenses					
			Sterilization of health care products General					
			requirements for characterization of a sterilizing					
			agent and the development, validation and routine					
ISO 14937	2009-10	N	control of a sterilization process for medical devices					
			Implants for surgery Two-part addition-cure					
ISO 14949	2001-10	N	silicone elastomers					
			Medical devices Application of risk management					
ISO 14971	2007-03	N	to medical devices	Р	N	JIS T 14971:2012	01.03.2012	See# 3 (802) on sheet1
			Sterile obturators for single use with over-needle					
ISO 14972	1998-12	N	peripheral intravascular catheters	Р	N	JIS T 3259: 2012	2012/10/1	392
			Anaesthetic and respiratory equipment					
ISO 15001	2010-06	N	Compatibility with oxygen					
100 45000	0000.07	N.	Flow-metering devices for connection to terminal					
ISO 15002	2008-07	N	units of medical gas pipeline systems		+			
			Ophthalmic instruments Fundamental requirements and test methods Part_1: General					
			requirements and test methods Part_1: General requirements applicable to all ophthalmic					
ISO 15004-1	2006-06	N	instruments					
100071	2000 00	13	Ophthalmic instruments - Fundamental					
			requirements and test methods - Part 2: Light					
ISO 15004-2	2007-02	N	hazard protection					
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			Disposable hanging devices for transfusion and					
ISO 15010	1998-06	N	infusion bottles Requirements and test methods					
ISO 15032	2000-04	N N	Prostheses Structural testing of hip units Dental elevators Part_1: General requirements					
ISO 15087-1	1999-11		·					
ISO 15087-2	2000-04	N	Dental elevators Part_2: Warwick James elevators					
ISO 15087-3	2000-05	N	Dental elevators Part_3: Cryer elevators					
ISO 15087-4	2000-05	N	Dental elevators Part_4: Coupland elevators					
ISO 15087-5	2000-05	N	Dental elevators Part_5: Bein elevators					
ISO 15087-6	2000-05	N	Dental elevators Part_6: Flohr elevators					
ISO 15098-1	1999-10	N	Dental tweezers Part_1: General requirements					
ISO 15098-2	2000-02	N	Dental tweezers Part_2: Meriam types					
ISO 15098-3	2000-02	N	Dental tweezers Part_3: College types					
			Self-adhesive hanging devices for infusion bottles					
			and injection vials Requirements and test					
ISO 15137	2005-07	N	methods					
			Implants for surgery Metal intramedullary nailing					
ISO 15142-1	2003-08	N	systems Part_1: Intramedullary nails					
			Implants for surgery Metal intramedullary nailing					
ISO 15142-2	2003-08	N	systems Part_2: Locking components					
			Implants for surgery Metal intramedullary nailing					
			systems Part_3: Connection devices and reamer					
ISO 15142-3	2003-08	N	diameter measurements					
			In vitro diagnostia modical devises Massurament					
			In vitro diagnostic medical devices Measurement of quantities in samples of biological origin					
			Requirements for content and presentation of					
ISO 15193	2009-05	N	reference measurement procedures					
100 10100	2000 00	.,	Total and Madadia Managara					
			In vitro diagnostic medical devices Measurement					
			of quantities in samples of biological origin					
			Requirements for certified reference materials and					
ISO 15194	2009-05	N	the content of supporting documentation					
	2013-05		In vitro diagnostic test systems Requirements for blood- glucose monitoring systems for self-testing in managing					
ISO 15197	2003-05	N	diabetes mellitus	Υ	N			AP19
100 10101		14	Clinical laboratory medicine - In vitro diagnostic	<u> </u>	N			Ai 10
			medical devices - Validation of user quality control					
ISO 15198	2004-07	N	procedures by the manufacturer					
100 10100	200101	.,	Medical devices - Symbols to be used with medical					
			device labels, labelling and information to be					
ISO 15223-1	2007-04	N	supplied Part_1: General requirements					
			Medical devices Symbols to be used with medical					
			device labels, labelling and information to be					
			supplied Part_1: General requirements;					
ISO 15223-1 AMD 1	2008-06	N	Amendment_1			1		
			Medical devices Symbols to be used with medical		1			
			device labels, labelling, and information to be					
ISO 15223-2	2010-01	NI NI	supplied Part_2: Symbol development, selection and validation					
100 10223-2	2010-01	N	anu vanuation		l		L	

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			Medical devices Quality management Medical					
ISO 15225	2010-05	N	device nomenclature data structure Ophthalmic optics and instruments Optical devices for					
ISO 15253	2000-09	N	enhancing low vision					
130 13233	2000-09	IN	Ophthalmic optics and instruments Electro-optical					
ISO 15254	2009-07	N	devices for enhancing low vision					
130 13234	2009-07	IN	Implants for surgery Requirements for production					
ISO 15374	1998-08	N	of forgings					
100 10074	1990-00	IN	or rorgings					
			Medical infusion bottles - Suspension devices for					
ISO 15375	2010-06	N	multiple use Requirements and test methods					
			Primary packaging materials for medicinal products					
			Particular requirements for the application of					
			ISO_9001:2008, with reference to Good Manufacturing Practice_(GMP)					
ISO 15378	2011-11	N	Practice_(GMP)					
ISO 15606	1999-12	N	Dental handpieces Air-powered scalers and scaler tips	Р	N	JIS T 5910:2005	25.03.2005	158
100 13000	1333-12	- 14	Derital Haridpieces_ All-powered scalers and scaler tips		11	010 1 00 10.2000		100
ISO 15621	2011-02	N	Urine-absorbing aids General guidelines on evaluation	Υ	N			AP36
ISO 1563	1990-09	N	Dental alginate impression material	Р	N	JIS T 6505:2005	25.03.2005	299
ISO 1564	1995-11	N	Dental aqueous impression materials based on agar	Р	N	JIS T 6512:2005	25.03.2005	301
			Cardiovascular implants and artificial organs					
			Hard-shell cardiotomy/venous reservoir systems					
ISO 15674	2009-04	N	(with/without filter) and soft venous reservoir bags	Υ	N			AP38
			Cardiovascular implants and artificial organs					
			Cardiopulmonary bypass systems Arterial blood					
ISO 15675	2009-04	N	line filters	Р	N	JIS T 3232:2011	29.07.2011	112 721
			Cardiovascular implants and artificial organs					
			Requirements for single-use tubing packs for					
			cardiopulmonary bypass and extracorporeal					
ISO 15676	2005-07	N	membrane oxygenation (ECMO)	Y	N			AP36
ISO 15747	2010-04	N	Plastic containers for intravenous injections					
			Ophthalmic instruments Endoilluminators					
ISO 15752	2010-01	N	Fundamental requirements and test methods for optical radiation safety					
130 13732	2010-01	IN	optical radiation salety					
			Medical infusion equipment Plastics caps with					
			inserted elastomeric liner for containers					
ISO 15759	2005-04	N	manufactured by the blow-fill-seal (BFS) process					
100 10100			Ophthalmic implants Ophthalmic viscosurgical					
ISO 15798	2010-01	N	devices					
			Implants for surgery Copolymers and blends					
ISO 15814	1999-11	N	based on polylactide In vitro degradation testing					
ISO 15841	2006-10	N	Dentistry Wires for use in orthodontics	Р	N	JIS T 6530:2009	25.06.2009	161
ISO 15854	2005-07	N	Dentistry Casting and baseplate waxes					
			Sterilization of health care products Chemical					
			indicators Guidance for selection, use and					
ISO 15882	2008-09	N	interpretation of results					
			Washer-disinfectors Part_1: General					
ISO 15883-1	2006-04	N	requirements, terms and definitions and tests		1			

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			Washer-disinfectors Part_2: Requirements and		
			tests for washer-disinfectors employing thermal		
			disinfection for surgical instruments, anaesthetic		
			equipment, bowls, dishes, receivers, utensils,		
ISO 15883-2	2006-04	N	glassware, etc.		
			Washer-disinfectors Part_3: Requirements and		
			tests for washer-disinfectors employing thermal		
ISO 15883-3	2006-04	N	disinfection for human waste containers		
			Washer-disinfectors Part_4: Requirements and		
			tests for washer-disinfectors employing chemical		
ISO 15883-4	2008-05	N	disinfection for thermolabile endoscopes		
			Washer-disinfectors Part_6: Requirements and		
			tests for washer-disinfectors employing thermal		
			disinfection for non-invasive, non-critical medical		
ISO 15883-6	2011-04	N	devices and healthcare equipment		
			Dentistry Casting investments and refractory die		
ISO 15912	2006-10	N	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 1	2011-07	N	materials		
100 100 12 / 11115 1	2011 01	.,	Urine-absorbing aids Basic principles for evaluation of		
			single-use adult-incontinence-absorbing aids from the		
ISO 16021	2000-11	N	perspective of users and caregivers		
			Ophthalmic optics Specifications for single-vision		
ISO 16034	2002-02	N	ready-to-wear near-vision spectacles		
			Ophthalmic optics Specifications for single-vision		
			ready-to-wear near- vision spectacles; Technical		
ISO 16034 Technical	Corri 2006-08	N	Corrigendum_1		
			Rubber condoms for clinical trials Measurement of		
ISO 16037	2002-05	N	physical properties		
			Rubber condoms for clinical trials Measurement of		
ISO 16037 AMD 1	2011-02	N	physical properties; Amendment_1		
			Rubber condoms Guidance on the use of ISO_4074 in		
ISO 16038	2005-11	N	the quality management of natural rubber latex condoms		
100 10030	2003-11	11	Implants for surgery - Minimum data sets for		
ISO 16054	2000-12	N	surgical implants		
100 10034	2000-12	IN	Dentistry Required elements for codification used in		
ISO 16059	2007-08	N	data exchange		
			Instrumentation for use in association with non-		
ISO 16061	2008-12	N	active surgical implants General requirements		
			Technical aids for persons with disability		
ISO 16201	2006-10	N	Environmental control systems for daily living		
			Ophthalmic optics - Information interchange for		
ISO 16284	2006-03	N	ophthalmic optical equipment		
			Aids for ostomy and incontinence Irrigation sets		
ISO 16391	2002-10	N	Requirements and test methods		
			Implants for surgery Acrylic resin cement		
			Flexural fatigue testing of acrylic resin cements		
ISO 16402	2008-05	N	used in orthopaedics		
ISO 16408	2004-04	N	Dentistry Oral hygiene products Oral rinses		

			Dentistry Oral hygiene products Manual interdental		
ISO 16409	2006-10	N	brushes		
			Dentistry Oral hygiene products Manual interdental		
ISO 16409 AMD 1	2010-02	N	brushes; Amendment_1		
			Implants for surgery Test solutions and		
			environmental conditions for static and dynamic		
			corrosion tests on implantable materials and		
ISO 16428	2005-04	N	medical devices		
			Implants for surgery Measurements of open-		
			circuit potential to assess corrosion behaviour of		
			metallic implantable materials and medical devices		
ISO 16429	2004-07	N	over extended time periods		
ISO 16628	2008-11	N	Tracheobronchial tubes - Sizing and marking		
			Ophthalmic implants Irrigating solutions for		
ISO 16671	2003-05	N	ophthalmic surgery		
ISO 16672	2003-02	N	Ophthalmic implants Ocular endotamponades		
			Wheelchair seating Part_1: Vocabulary, reference axis		
			convention and measures for body segments, posture		
ISO 16840-1	2006-03	N	and postural support surfaces		
			Wheelchair seating Part_2: Determination of physical		
			and mechanical characteristics of devices intended to		
ISO 16840-2	2007-07	N	manage tissue integrity Seat cushions		
			Wheelchair seating Part_3: Determination of static,		
			impact and repetitive load strengths for postural support		
ISO 16840-3	2006-07	N	devices		
			Wheelchair seating Part_4: Seating systems for use in		
ISO 16840-4	2009-03	N	motor vehicles		
			Health informatics Public key infrastructure Part_1:		
ISO 17090-1	2008-02	N	Overview of digital certificate services		
			Health informatics Public key infrastructure Part_2: Certificate profile		
ISO 17090-2	2008-02	N	Health informatics Public key infrastructure Part_3:		
ISO 17090-3	2008-02	N	Policy management of certification authority		
150 17090-3	2008-02	N	Health informatics Vocabulary for terminological		
ISO 17115	2007-07	N	systems		
130 17113	2007-07	IN	Urine-absorbing aids for incontinence - Test methods for		
			characterizing polymer-based absorbent materials		
ISO 17190-1	2001-12	N	Part 1: Determination of pH		
100 17 100 1	2001 12	''	Urine-absorbing aids for incontinence Test methods for		
			characterizing polymer-based absorbent materials		
			Part_10: Determination of extractable polymer content by		
ISO 17190-10	2001-12	N	potentiometric titration		
			Urine-absorbing aids for incontinence Test methods for		
			characterizing polymer-based absorbent materials		
ISO 17190-11	2001-12	N	Part_11: Determination of content of respirable particles		
			Urine-absorbing aids for incontinence Test methods for		
			characterizing polymer-based absorbent materials		
ISO 17190-2	2001-12	N	Part_2: Determination of amount of residual monomers		
			Urine absorbing aids for incontinence Test methods for		
			characterizing polymer-based absorbent materials Part_3: Determination of particle size distribution by sieve		
100 17100 2	2004 42	N.	fractionation		
ISO 17190-3	2001-12	N	nactionation		

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			Urine-absorbing aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials					
			Part_4: Determination of moisture content by mass loss upon heating					
ISO 17190-4	2001-12	N	Urine-absorbing aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials -					
			Part 5: Gravimetric determination of free swell capacity in					
ISO 17190-5	2001-12	N	saline solution					
100 17 100 0	2001 12	.,	Urine-absorping aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials					
			Part_6: Gravimetric determination of fluid retention					
ISO 17190-6	2001-12	N	capacity in saline solution after centrifugation					
			Urine-absorbing aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials					
			Part_7: Gravimetric determination of absorption under pressure					
ISO 17190-7	2001-12	N						
			Urine-absorping aids for incontinence Test methods for characterizing polymer-based absorbent materials					
ISO 17190-8	2001-12	N	Part 8: Gravimetric determination of flowrate					
130 17 190-6	2001-12	IN	Urine-absorbing aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials					
ISO 17190-9	2001-12	N	Part_9: Gravimetric determination of density					
			Urine-absorbing aids for incontinence Test methods for					
			characterizing polymer-based absorbent materials					
			Part_9: Gravimetric determination of density; Technical					
ISO 17190-9 Technical (Corrig 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					
130 17 191	2004-02	IN .	Health informatics - Messages and communication -					
ISO 17432	2004-12	N	Web access to DICOM persistent objects					
			Sleep apnoea breathing therapy - Part 1: Sleep					
ISO 17510-1	2007-10	N	apnoea breathing therapy equipment					
			Sleep apnoea breathing therapy Part_2: Masks					
ISO 17510-2	2007-10	N	and application 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 of oral anticoagulant					
ISO 17593	2007-04	N	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					
		1	Part_1: Requirements for the development,					NALILIAN NASSISIAN SI LINI SI
		1	validation and routine control of a sterilization					MHLW Ministerial Notice
ISO 17665-1	2006-08	N	process for medical devices	Υ	N	JIS T 0816-1:2010	25.02.2010	AP42
			Wear of implant materials Polymer and metal					
ISO 17853	2011-03	N	wear particles Isolation and characterization					
100 1707 1	0044.00	l	Dentistry Shanks for rotary instruments Part_1: Shanks made of metals					
ISO 1797-1	2011-08	N	Sharks hidde of filetals					

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ISO 1797-2	1992-02	N	Dental rotary instruments; shanks; part_2: shanks made of plastics				
ISO 18084	2011-09	N	Press tools for tablets Punches and dies				
			Health informatics Integration of a reference				
ISO 18104	2003-12	N	terminology model for nursing				
			In vitro diagnostic medical devices Information				
			supplied by the manufacturer (labelling) Part_1:				
ISO 18113-1	2009-12	N	Terms, definitions and general requirements				
			In vitre diagnostic modical devices. Information				
			In vitro diagnostic medical devices Information supplied by the manufacturer (labelling) Part_2:				
ISO 18113-2	2009-12	N	In vitro diagnostic reagents for professional use				
100 10113-2	2003-12	IN IN	in vito diagnostic reagents for professional use				
			In vitro diagnostic medical devices Information				
			supplied by the manufacturer (labelling) Part_3:				
ISO 18113-3	2009-12	N	In vitro diagnostic instruments for professional use				
			In vitro diagnostic medical devices Information				
			supplied by the manufacturer (labelling) Part_4:				
ISO 18113-4	2009-12	N	In vitro diagnostic reagents for self-testing				
			In vitro diagnostic medical devices Information				
100 40440 5	0000.40		supplied by the manufacturer (labelling) Part_5:				
ISO 18113-5	2009-12	N	In vitro diagnostic instruments for self-testing				
			In vitro diagnostic medical devices Measurement				
			of quantities in biological samples - Metrological				
			traceability of values for catalytic concentration of				
ISO 18153	2003-08	N	enzymes assigned calibrators and control materials				
			Implants for surgery Wear of total intervertebral				
			spinal disc prostheses Part_1: Loading and				
			displacement parameters for wear testing and				
ISO 18192-1	2011-03	N	corresponding environmental conditions for test				
			Implants for surgery Wear of total intervertebral				
ISO 18192-2	2010-06	N	spinal disc prostheses Part_2: Nucleus replacements				
150 18192-2	2010-06	IN	replacements				
			Health Informatics Messages and communication				
ISO 18232	2006-04	N	Format of length limited globally unique string identifiers				
			Health informatics Requirements for an electronic				
ISO 18308	2011-04	N	health record architecture				
			Ophthalmic optics Contact lenses Part_1:				
100 40000 4	0000 00		Vocabulary, classification system and				
ISO 18369-1	2006-08	N	recommendations for labelling specifications			_	
			Ophthalmic optics Contact lenses Part_1: Vocabulary, classification system and				
			recommendations for labelling specifications;				
ISO 18369-1 AMD 1	2009-02	N	Amendment 1				
		<u> </u>	Ophthalmic optics Contact lenses Part_2:				
ISO 18369-2	2006-08	N	Tolerances				
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		T	Onbthalmia antica Contact langua Bort 2:					
100 10360 3	2006.09	N.	Ophthalmic optics Contact lenses Part_3:	Υ	N			A D4
ISO 18369-3	2006-08	N	Measurement methods Ophthalmic optics Contact lenses Part_4:	r	IN			AP1
			Physicochemical properties of contact lens					
ISO 18369-4	2006-08	N	materials	Y	N			AP1
100 10003-4	2000-00	IN	Sterilization of health care products - Biological	ī	IN			Ar I
ISO 18472	2006-06	N	and chemical indicators Test equipment					
100 10472	2000-00	IN	Transportable liquid oxygen systems for medical		+			
ISO 18777	2005-02	l N	use - Particular requirements					
130 16777	2003-02	IN	Respiratory equipment Infant monitors					
ISO 18778	2005-02	N	Particular requirements					
100 10770	2003-02	IN	Medical devices for conserving oxygen and oxygen					
ISO 18779	2005-02	N	mixtures Particular requirements					
100 10770	2000 02		Health informatics Clinical analyser interfaces to					
ISO 18812	2003-03	N	laboratory information systems Use profiles					
			In vitro diagnostic medical devices - Information					
1			supplied by the manufacturer with in vitro					
ISO 19001	2002-11	N	diagnostic reagents for staining in biology					
ISO 19054	2005-07	N	Rail systems for supporting medical equipment					
ISO 1942	2009-12	N	Dentistry Vocabulary					
ISO 19980	2005-08	N	Ophthalmic instruments Corneal topographers					
			Aerosol drug delivery device design verification -					
ISO 20072	2009-08	N						
			Requirements and test methods Dentistry Manual toothbrushes General requirements					
ISO 20126	2012-01	N	and test methods					
			Dentistry Powered toothbrushes General					
ISO 20127	2005-03	N	requirements and test methods					
			Implants for surgery Metallic materials					
			Classification of microstructures for alpha+beta					
ISO 20160	2006-05	N	titanium alloy bars					
ISO 20301	2006-11	N	Health informatics Health cards General characteristics					
150 20301	2006-11	IN .	Health informatics Health cards Numbering system					
ISO 20302	2006-12	N	and registration procedure for issuer identifiers					
130 20302	2000-12	IN	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					
ISO 20776-1	2006-11	N	aerobic bacteria involved in infectious diseases					
100 20770 1	2000 11	i i	Clinical laboratory testing and in vitro diagnostic					
			test systems Susceptibility testing of infectious					
			agents and evaluation of performance of					
			antimicrobial susceptibility test devices - Part 2:					
			Evaluation of performance of antimicrobial					
ISO 20776-2	2007-07	N	susceptibility test devices					
- /		1	Dentistry Base polymers Part_1: Denture base					
ISO 20795-1	2008-08	N	polymers	Р	N	JIS T 6501:2012	01.03.2012	242· AP36
			Dentistry Base polymers Part_1: Denture base					
ISO 20795-1 Technical	Corrig 2009-02	N	polymers; Technical Corrigendum_1	Р	N	JIS T 6501:2012	01.03.2012	242
			Dentistry Base polymers Part_2: Orthodontic base					
ISO 20795-2	2010-03	N	polymers					

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			Sterilization of health care products Dry heat					
			Requirements for the development, validation and					
			routine control of a sterilization process for medical					
ISO 20857	2010-08	N	devices					
			Health informatics Harmonized data types for					
ISO 21090	2011-02	N	information interchange					
			Medical gloves Determination of removable					
ISO 21171	2006-05	N	surface powder					
100 04500	0004.00		Dentistry Materials used for dental equipment surfaces Determination of resistance to chemical disinfectants					
ISO 21530	2004-06	N						
ISO 21531	2009-02	N	Dentistry Graphical symbols for dental instruments					
			Dentistry Reusable cartridge syringes intended for					
ISO 21533	2003-06	N	intraligamentary injections					
ISO 21533 Technical Co	2000 12	N	Dentistry Reusable cartridge syringes intended for intraligamentary injections; Technical Corrigendum_1					
150 21533 Technical Co	ornger 2009-12	IN						
100 04504	0007.40		Non-active surgical implants Joint replacement					
ISO 21534	2007-10	N	implants Particular requirements					
			Non-active surgical implants Joint replacement					
100 04505	0007.40	N.	implants Specific requirements for hip-joint	Υ	N			01.4
ISO 21535	2007-10	N	replacement implants	Y	N N			GL1
			Non-active surgical implants Joint replacement					
100 04500	0007.40	N.	implants Specific requirements for knee-joint replacement implants					
ISO 21536	2007-10	N	Health informatics - Patient healthcard data - Part 1:					
ISO 21549-1	2004-05	N	General structure					
100 210 10 1	2004 00		Health informatics - Patient healthcard data - Part 2:					
ISO 21549-2	2004-05	N	Common objects					
			Health informatics Patient healthcard data Part_3:					
ISO 21549-3	2004-05	N	Limited clinical data					
			Health informatics Patient healthcard data Part_4:					
ISO 21549-4	2006-11	N	Extended clinical data					
			Health informatics Patient healthcard data Part_5:					
ISO 21549-5	2008-04	N	Identification data					
100 04540 0	0000.04	N.	Health informatics Patient healthcard data Part_6: Administrative data					
ISO 21549-6	2008-04	N	Health informatics Patient healthcard data Part_7:					
ISO 21549-7	2007-06	N	Medication data					
130 21343-7	2007-00	IN	Health informatics - Patient healthcard data - Part 8:					
ISO 21549-8	2010-06	N	Links					
100 210 10 0	2010 00	.,	Dental rotary instruments; nominal diameters and					
ISO 2157	1992-06	N	designation code number					
ISO 21606	2007-06	N	Dentistry Elastomeric auxiliaries for use in orthodontics	Р	N	JIS T 6531:2012	01.03.2012	173
			Needle-free injectors for medical use					
ISO 21649	2006-06	N	Requirements and test methods					
			Health informatics Health indicators conceptual framework					
ISO 21667	2010-12	N						
ISO 21671	2006-07	N	Dentistry Rotary polishers					
ISO 21671 AMD 1	2011-04	N	Dentistry Rotary polishers; Amendment_1					
			Dentistry Periodontal probes Part_1: General					
ISO 21672-1	2012-04	N	requirements					
		1	High-pressure flexible connections for use with					
ISO 21969	2009-10	N	medical gas systems					

ISO 21987	2009-10	N	Onbtholmic entire Mounted an estacle langer				1	
			Ophthalmic optics Mounted spectacle lenses Dentistry Artificial teeth for dental prostheses					
ISO 22112	2005-11	N	Dentistry Artificial teeth for dental prostneses Dentistry Manual toothbrushes Resistance of tufted					
100 00054	0005.00		portion to deflection					
ISO 22254	2005-08	N	Dentistry Dental handpieces Electrical-powered					
ISO 22374	2005-09	N	scalers and scaler tips	Р	N	JIS T 5911:2012	01.03.2012	157
150 22374	2005-09	IN	Transfer sets for pharmaceutical preparations	F	IN	JIS 1 5911.2012	01.03.2012	157
ISO 22413	2010-06	N	Requirements and test methods					
150 22413	2010-06	IN	· · · · · · · · · · · · · · · · · · ·					
			Medical devices utilizing animal tissues and their					
			derivatives Part_1: Application of risk					
ISO 22442-1	2007-12	N	management					
			Medical devices utilizing animal tissues and their					
			derivatives Part_2: Controls on sourcing,					
ISO 22442-2	2007-12	N	collection and handling					
			Medical devices utilizing animal tissues and their					
			derivatives Part_3: Validation of the elimination					
			and/or inactivation of viruses and transmissible					
ISO 22442-3	2007-12	N	spongiform encephalopathy (TSE) agents					
			External limb prostheses and external orthoses					
ISO 22523	2006-10	N	Requirements and test methods					
			Clothing for protection against infectious agents Medical					
			face masks Test method for resistance against					
100 0000			penetration by synthetic blood (fixed volume, horizontally					
ISO 22609	2004-12	N	projected)					
			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 dry microbial					
ISO 22612	2005-03	N	penetration					
						UO T 0445 0040	0040/0/4	203
						JIS T 6115:2013	2013/3/1	185,AP36
						JIS T 6116:2012 JIS T 6118:2012	2012/7/1 2012/7/1	188.AP36
						JIS T 6121:2013	2012///1	214
			Dentistry Metallic materials for fixed and removable			JIS T 6121:2013	2013/3/1	186
ISO 22674	2006-11	N	restorations and appliances	P	N	JIS T 6123:2013	2013/3/1	209
100 22014	2000 11	1,	Prosthetics Testing of ankle-foot devices and foot	•	- ''	0.0 1 0120.2010	2010/0/1	255
ISO 22675	2006-10	N	units - Requirements and test methods					
ISO 22715	2006-04	N	Cosmetics Packaging and labelling					
130 227 13	2000-04	IN	Cosmetics Good Manufacturing Practices (GMP)					
ISO 22716	2007-11	N	Guidelines on Good Manufacturing Practices					
130 227 10	2007-11	IN	Dentistry Implantable materials for bone filling and					
			augmentation in oral and maxillofacial surgery Contents					
ISO 22794	2007-07	N	of a technical file					
		1	Dentistry Membrane materials for guided tissue			1		
			regeneration in oral and maxillofacial surgery Contents					
ISO 22803	2004-09	N	of a technical file					
			Health informatics Guidelines on data protection to					
ISO 22857	2004-04	N	facilitate trans-border flows of personal health information					
			Implants for surgery In vitro evaluation for apatite-					
ISO 23317	2007-06	N	forming ability of implant materials					
			. • • • • • • • • • • • • • • • • • • •					

		1	Describing and the Character of the state of				1	
			Breathing system filters for anaesthetic and					
			respiratory use Part_1: Salt test method to					
ISO 23328-1	2003-08	N	assess filtration performance	Y	N	JIS T 7211:2005	25.03.2005	539 · 543 · 540 · 113
100 00000 0	0000 10		Breathing system filters for anaesthetic and				05 00 0005	500 540 540 440
ISO 23328-2	2002-10	N	respiratory use Part_2: Non-filtration aspects Male condoms - Requirements and test methods for	Y	N	JIS T 7212:2005	25.03.2005	539 · 543 · 540 · 113
ISO 23409	2011-02	N	condoms made from synthetic materials					
130 23409	2011-02	IN	Guidance for the preparation and quality					
			management of fluids for haemodialysis and					
ISO 23500	2011-05	N	related therapies					
100 20000	2011 00	.,	Assistive products for blind and vision-impaired persons					
ISO 23599	2012-03	N	Tactile walking surface indicators					
			Assistive products for persons with vision impairments					
ISO 23600	2007-11	N	and persons with vision and hearing impairments Acoustic and tactile signals for pedestrian traffic lights					
130 23000	2007-11	IN	In vitro diagnostic medical devices - Evaluation of					
ISO 23640	2011-12	N	stability of in vitro diagnostic reagents					
20040	2011-12	IN	Anaesthetic and respiratory equipment Peak		+			
			expiratory flow meters for the assessment of					
			pulmonary function in spontaneously breathing					
ISO 23747	2007-07	N	humans					
			Sharps injury protection Requirements and test					
			methods Sharps protection features for single-					
			use hypodermic needles, introducers for catheters					
ISO 23908	2011-06	N	and needles used for blood sampling					
			Ophthalmic optics and instruments Reporting					
ISO 24157	2008-07	N	aberrations of the human eye Skin barrier for ostomy aids Vocabulary					
ISO 24214	2006-11	N	· - ·					211 212
ISO 24234	2004-10	N	Dentistry Mercury and alloys for dental amalgam Dentistry Mercury and alloys for dental amalgam	Р	N	JIS T 6127:2008	25.11.2008	211-212
			Amendment_1: Requirements for marking and					
ISO 24234 AMD 1	2011-08	N	manufacturer's instructions concerning mercury					
			Tips for assistive products for walking Requirements					
ISO 24415-1	2009-04	N	and test methods Part_1: Friction of tips					
			Tips for assistive products for walking Requirements					
ISO 24415-2	2011-08	N	and test methods - Part 2: Durability of tips for crutches					
100 24410 2	2011 00		Ergonomics Accessible design Auditory signals for					
ISO 24500	2010-10	N	consumer products					
			Ergonomics Accessible design Sound pressure levels					
ISO 24501	2010-12	N	of auditory signals for consumer products					
100 04500	2010.10		Ergonomics Accessible design Specification of age- related luminance contrast for coloured light					
ISO 24502	2010-12	N	Ergonomics Accessible design Tactile dots and bars					
ISO 24503	2011-01	N	on consumer products					
	20		<u> </u>					
			Sterilization of medical devices Low temperature					
			steam and formaldehyde Requirements for					
			development, validation and routine control of a		1			
ISO 25424	2009-09	N	sterilization process for medical devices					
			Cardiovascular implants Endovascular devices					
ISO 25539-1	2003-03	N	Part_1: Endovascular prostheses					

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			Cardiovascular implants Endovascular devices	ĺ			
			Part_1: Endovascular prostheses; Amendment_1:	ĺ			
ISO 25539-1 AMD 1	2005-07	N	Test methods				
			Cardiovascular implants Endovascular devices	ĺ			
ISO 25539-2	2008-09	N	Part_2: Vascular stents				
			Cardiovascular implants Endovascular devices	ĺ			
ISO 25539-3	2011-12	N	Part_3: Vena cava filters Health informatics Genomic Sequence Variation				
ISO 25720	2009-08	N	Markup Language (GSVML)	ĺ			
	2011-07	N	Female condoms Requirements and test methods	 			
ISO 25841	2011-07	IN .	_ ·				
100 00700	0000 04	N.	Water treatment equipment for haemodialysis	ĺ			
ISO 26722	2009-04	N	applications and related therapies				
			Anaesthetic and respiratory equipment Spirometers intended for the measurement of time	ĺ			
ISO 26782	2009-07	l N	forced expired volumes in humans	ĺ			
130 20702	2009-07	IN	Anaesthetic and respiratory equipment -				
			Spirometers intended for the measurement of time	ĺ			
			forced expired volumes in humans; Technical	ĺ			
ISO 26782 Technical C	orri 2009-11	N	Corrigendum 1	ĺ			
100 Z070Z TCCIIIICAI O	01112003 11	11	Anaesthetic and respiratory equipment User-				
			applied labels for syringes containing drugs used	ĺ			
			during anaesthesia Colours, design and	ĺ			
ISO 26825	2008-08	N	performance	ĺ			
ISO 27020	2010-12	N	Dentistry Brackets and tubes for use in orthodontics				
			Cardiac rhythm management devices Symbols to				
			be used with cardiac rhythm management device	ĺ			
			labels, and information to be supplied - General	ĺ			
ISO 27185	2012-02	N	requirements	ĺ			
			Active implantable medical devices Four-pole				
			connector system for implantable cardiac rhythm	ĺ			
			management devices Dimensional and test	ĺ			
ISO 27186	2010-03	N	requirements	ĺ			
			Anaesthetic and respiratory equipment				
ISO 27427	2010-03	N	Nebulizing systems and components				
			Health informatics Information security management in	ĺ			
ISO 27799	2008-07	N	health using ISO/IEC_2702				
ISO 28158	2010-07	N	Dentistry Integrated dental floss and handles				
ISO 28319	2010-05	N	Dentistry Laser welding				
ISO 28399	2011-01	N	Dentistry Products for external tooth bleaching				
			Medical devices Non-electrically driven portable				
ISO 28620	2010-02	N	infusion devices				
			Nanotechnologies Endotoxin test on				
			nanomaterial samples for in vitro systems	ĺ			
ISO 29701	2010-09	N	Limulus amebocyte lysate (LAL) test				
			Prostheses and orthoses Factors to be included when	ĺ			
			describing physical activity of a person who has had a	ĺ			
ISO 29781	2008-12	N	lower limb amputation(s) or who has a deficiency of a lower limb segment(s) present at birth	1			
130 29/01	2000-12	IN	Prostheses and orthoses - Factors to be considered	<u> </u>			
			when specifying a prosthesis for a person who has had a	Í			
ISO 29782	2008-12	N	lower limb amputation	1			
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r		1	<u> </u>		1			
			Prosthetics and orthotics Vocabulary Part_1:					
ISO 29783-1	2008-12	N	Normal gait Condoms - Determination of nitrosamines migrating from		ļ			
ISO 29941	2010-12	N	natural rubber latex condoms					
ISO 29942	2011-07	N	Prophylactic dams Requirements and test methods					
			Dentistry Zinc oxide/eugenol cements and zinc					
ISO 3107	2011-03	N	oxide/non-eugenol cements	Р	N	JIS T 6610: 2013	01.09.2013	256 · 257 · 291
			Gas cylinders for medical use; Marking for					
ISO 32	1977-05	N	identification of content					
ISO 3630-1	2008-02	N	Dentistry Root-canal instruments Part_1: General requirements and test methods					
ISO 3630-2	2000-12	N	Dental root-canal instruments Part_2: Enlargers					
			Dental root-canal instruments; part_3: condensers,					
ISO 3630-3	1994-03	N	pluggers and spreaders					
			Dentistry Root canal instruments Part_4: Auxiliary					
ISO 3630-4	2009-07	N	instruments Dentistry Endodontic instruments Part_5: Shaping					
ISO 3630-5	2011-10	N	and cleaning instruments					
100 0000 0	2011 10	.,	Dental rotary instruments Burs Part_1: Steel and					
ISO 3823-1	1997-08	N	carbide burs					
			Dentistry Rotary bur instruments Part_2: Finishing					
ISO 3823-2	2003-05	N	burs					
ISO 3823-2 AMD 1	2008-07	N	Dentistry Rotary bur instruments Part_2: Finishing burs; Amendment_1					
ISO 3823-2 AMD 1	2008-07	IN .	Plastics collapsible containers for human blood and					
			blood components Part_1: Conventional					
ISO 3826-1	2003-11	N	containers	Р	N	JIS T 3217:2011	29.07.2011	98 · 636 · 676 · 677
100 0020 1	2000 11	- ''	Plastics collapsible containers for human blood and	· · · · · · · · · · · · · · · · · · ·	- ''	010 1 0217.2011	20.07.2011	00 000 010 011
			blood components Part_2: Graphical symbols for					
ISO 3826-2	2008-08	N	use on labels and instruction leaflets					
			Plastics collapsible containers for human blood and					
			blood components Part_3: Blood bag systems					
ISO 3826-3	2006-09	N	with integrated features					
			Acoustics Reference zero for the calibration of					
			audiometric equipment Part_1: Reference					
100 000 4	4000 44	N.	equivalent threshold sound pressure levels for pure					
ISO 389-1	1998-11	N	tones and supra-aural earphones Acoustics - Reference zero for the calibration of					
			audiometric equipment Part_2: Reference					
			equivalent threshold sound pressure levels for pure					
ISO 389-2	1994-07	N	tones and insert earphones					
	100101		Acoustics Reference zero for the calibration of					
			audiometric equipment Part_3: Reference					
			equivalent threshold force levels for pure tones and					
ISO 389-3	1994-10	N	bone vibrators					
			Acoustics Reference zero for the calibration of					
			audiometric equipment Part_3: Reference					
			equivalent treshold force levels for pure tones and					
ISO 389-3 Technical C	Corrig 1995-08	N	bone vibrators; Technical corrigendum_1					
			Acoustics Reference zero for the calibration of					
100 200 4	1994-10	N	audiometric equipment Part_4: Reference levels					
ISO 389-4	[1994-10	N	for narrow-band masking noise					

		1	IA .: D.(1			1
			Acoustics Reference zero for the calibration of					
			audiometric equipment Part_6: Reference					
ISO 389-6	2007-07	N	threshold of hearing for test signals of short duration					
130 303-0	2007-07	IN	Acoustics - Reference zero for the calibration of					
			audiometric equipment Part_7: Reference					
			threshold of hearing under free-field and diffuse-					
ISO 389-7	2005-11	N	field listening conditions					
100 303 7	2000 11	11	Acoustics Reference zero for the calibration of					
			audiometric equipment Part_8: Reference					
			equivalent threshold sound pressure levels for pure					
ISO 389-8	2004-05	N	tones and circumaural earphones					
			Acoustics - Reference zero for the calibration of					
			audiometric equipment Part_9: Preferred test					
			conditions for the determination of reference					
ISO 389-9	2009-05	N	hearing threshold levels					
			Dentistry Designation system for teeth and areas of the					
ISO 3950	2009-05	N	oral cavity					
ISO 3964	1982-12	N	Dental handpieces; Coupling dimensions					
						JIS T 6514: 2013	2013/3/1	266
ISO 4049	2009-10	N	Dentistry Polymer-based restorative materials	Р	N	JIS T 6523:2005	2005/3/25	280
			Dentistry Information system on the location of dental					
			equipment in the working area of the oral health care					
ISO 4073	2009-07	N	provider Natural latex rubber condoms Requirements and test					
ISO 4074	2002-02	N	methods					
130 4074	2002-02	IN	Natural latex rubber condoms - Requirements and test					
ISO 4074 Technical Co	orrigen 2003-10	N	methods; Technical Corrigendum_1					
			Natural latex rubber condoms Requirements and test					
ISO 4074 Technical Co	orrigend 2008-04	N	methods; Technical Corrigendum_2					
			Anaesthetic and respiratory equipment					
ISO 4135	2001-08	N	Vocabulary					
ISO 4823	2000-12	N	Dentistry Elastomeric impression materials	Р	N	JIS T 6513:2005	25.03.2005	300
			Dentistry Elastomeric impression materials;					
ISO 4823 AMD 1	2007-07	N	Amendment_1					
			Dentistry Elastomeric impression materials; Technical Corrigendum 1					
ISO 4823 Technical Co	orrigen(2004-07	N	Corngendum_1					04 05 00 440 445 440 5
								84 · 85 · 86 · 113 · 115 · 118 · 5
								10 · 524 · 539 · 540 · 542 · 543
			Anaesthetic and respiratory equipment Conical					· 544 · 547 · 548 · 653 · 768 · 7
ISO 5356-1	2004-05	N	connectors Part_1: Cones and sockets	Р	N	JIS T 7201-2-1:1999	30.04.1999	91 · 116 · 117 ·
			Anaesthetic and respiratory equipment Conical					
			connectors Part_2: Screw-threaded weight-					
ISO 5356-2	2006-09	N	bearing connectors	Р	N	JIS T 7201-2-2:1999	30.04.1999	539 540
ISO 5358	1992-01	N	Anaesthetic machines for use with humans	Р	N	JIS T 7201-1:1999	30.04.1999	766
	-		Low-pressure hose assemblies for use with					
ISO 5359	2008-06	N	medical gases	Р	N	JIS T 7111:2006	01.11.2006	524 · 765 · 767
			Low-pressure hose assemblies for use with	•	1			
ISO 5359 AMD 1	2011-12	N	medical gases; Amendment 1					
			Anaesthetic vaporizers Agent-specific filling					
ISO 5360	2012-01	N	systems		1			

			Anaesthetic and respiratory equipment Tracheal		1			
SO 5361	1999-09	N	tubes and connectors	Р	N	JIS T 7221:2011	29.07.2011	84 · 85
SO 5361-4	1987-12	N	Tracheal tubes; Part 4 : Cole type	P	N	JIS T 7224:1993	13.07.1993	84
SO 5362	2006-06	N	Anaesthetic reservoir bags	Р	N	JIS T 7201-3:2005	25.03.2005	118· 539· 540
			Anaesthetic and respiratory equipment					
SO 5364	2008-07	N	Oropharyngeal airways					
			Anaesthetic and respiratory equipment					
00 5000 4	2000 40		Tracheostomy tubes Part_1: Tubes and connectors for use in adults	_			00.07.0044	
SO 5366-1	2000-12	N	Anaesthetic and respiratory equipment	Р	N	JIS T 7227:2011	29.07.2011	86
			Tracheostomy tubes Part_3: Paediatric					
SO 5366-3	2001-08	N	tracheostomy tubes	Р	l N	JIS T 7227:2011	29.07.2011	86
			Anaesthetic and respiratory equipment					
			Tracheostomy tubes Part_3: Paediatric					
SO 5366-3 Technic	al Cort 2003-01	N	tracheostomy tubes; Technical Corrigendum_1					
			Breathing tubes intended for use with anaesthetic					
SO 5367	2000-06	N	apparatus and ventilators	Р	N	JIS T 7201-4:2005	25.03.2005	116 509 539 540 115
SO 5832-1	2007-06	N	Implants for surgery Metallic materials Part_1: Wrought stainless steel	Υ	N			AP28 AP36、GL4、5、7、8
SU 3832-1	2007-06	IN	vrought stainless steel	Ť	IN IN			AP36, GL4, 5, 7, 8
			Implants for surgery Metallic materials Part_1:					
SO 5832-1 Technic	al Corr 2008-04	N	Wrought stainless steel; Technical Corrigendum_1					
			Implants for surgery Metallic materials Part_11:					
SO 5832-11	1994-09	N	Wrought titanium 6-aluminium 7-niobium alloy	Υ	N			AP36、GL4、5、8
			Implants for surgery Metallic materials Part_12:					
SO 5832-12	2007-05	N	Wrought cobalt-chromium-molybdenum alloy					
			Implants for surgery Metallic materials Part_12:					
			Wrought cobalt-chromium-molybdenum alloy;					
SO 5832-12 Techn	ical Co 2008-09	N	Technical Corrigendum_1					
			Implants for surgery Metallic materials Part_14: Wrought titanium 15-molybdenum 5-zirconium 3-					
SO 5832-14	2007-10	N	aluminium alloy					
			Implants for surgery Metallic materials Part_2:					
SO 5832-2	1999-07	N	Unalloyed titanium	Υ	N			AP27 • GL4、5、7、8
SO 5832-3	1996-07	N	Implants for surgery Metallic materials Part_3: Wrought titanium 6-aluminium 4-vanadium alloy	Υ	N			AP27 • AP36 • GL4、5、7、8
30 3032-3	1990-07	IN	virought thanium 6-aluminium 4-variaulum alloy	T	IN			AF27-AF30-GL4, 5, 7, 8
			Implants for surgery Metallic materials Part_4:					
SO 5832-4	1996-07	N	Cobalt-chromium-molybdenum casting alloy					
00 5000 5	2005.40		Implants for surgery Metallic materials Part_5:					.=== .===
SO 5832-5	2005-10	N	Wrought cobalt-chromium-tungsten-nickel alloy	Y	N			AP27·AP28
			Implants for surgery Metallic materials Part_6:					
SO 5832-6	1997-07	N	Wrought cobalt-nickel-chromium-molybdenum alloy	Υ	N			AP27
			Implants for surgery; metallic materials; part_7:					
			forgeable and cold-formed cobalt-chromium-nickel-					
SO 5832-7	1994-02	N	molybdenum-iron alloy	Υ	N			AP27 • AP28

	ı	1			T		1
			Implants for surgery Metallic materials Part_8:				
			Wrought cobalt-nickel-chromium-molybdenum-				
ISO 5832-8	1997-07	N	tungsten-iron alloy	Y	N		AP27
100 5000 6	000= 00		Implants for surgery Metallic materials Part_9:				
ISO 5832-9	2007-06	N	Wrought high nitrogen stainless steel	Y	N		GL4
ISO 5833	2002-05	N	Implants for surgery Acrylic resin cements				
100 5004 4	0005.00	N.	Implants for surgery Ultra-high-molecular-weight				
ISO 5834-1	2005-06	N	polyethylene Part_1: Powder form Implants for surgery - Ultra-high-molecular-weight				
			polyethylene Part_1: Powder form; Technical				
ISO 5834-1 Technical	Cort 2007-05	N	Corrigendum_1				
100 3034-1 Technical	2007-03	IN	Implants for surgery Ultra-high-molecular-weight				
ISO 5834-2	2011-08	N	polyethylene Part_2: Moulded forms				
100 000+ 2	2011 00		Implants for surgery - Ultra-high-molecular-weight				
			polyethylene Part_3: Accelerated ageing				
ISO 5834-3	2005-07	N	methods				
			Implants for surgery Ultra-high-molecular-weight				
			polyethylene Part_4: Oxidation index				
ISO 5834-4	2005-05	N	measurement method				
			Implants for surgery Ultra-high-molecular-weight				
			polyethylene Part_5: Morphology assessment				
ISO 5834-5	2005-06	N	method				
			Implants for surgery; metal bone screws with				
			hexagonal drive connection, spherical under-				
ISO 5835	1991-01	N	surface of head, asymmetrical thread; dimensions				
			Implants for surgery; metal bone plates; holes				
			corresponding to screws with asymmetrical thread				
ISO 5836	1988-12	N	and spherical under-surface				
			Implants for surgery; Intramedullary nailing systems; Part 1 : Intramedullary nails with				
ISO 5837-1	1985-06	N	cloverleaf or V-shaped cross-section				
130 3037-1	1905-00	IN	Implants for surgery; Intramedullary nailing				
ISO 5837-2	1980-11	N	systems; Part 2 : Medullary pins				
100 3037-2	1300-11	IN	Implants for surgery Skeletal pins and wires			+	
ISO 5838-1	1995-11	N	Part_1: Material and mechanical requirements				
100 0000 1	1000 11	.,	Implants for surgery; skeletal pins and wires;				
ISO 5838-2	1991-01	N	part_2: Steinmann skeletal pins; dimensions				
			Implants for surgery; skeletal pins and wires;				
ISO 5838-3	1993-09	N	part_3: Kirschner skeletal wires				
			Cardiovascular implants Cardiac valve				
ISO 5840	2005-03	N	prostheses				
			Implants for surgery Cardiac pacemakers				
			Part_2: Reporting of clinical performance of				
ISO 5841-2	2000-10	N	populations of pulse generators or leads				
			Implants for surgery Cardiac pacemakers				
			Part_3: Low-profile connectors [IS-1] for				
ISO 5841-3	2000-10	N	implantable pacemakers				
			Implants for surgery Cardiac pacemakers				
			Part_3: Low-profile connectors (IS-1) for				
100 5044 0 7 / / /			implantable pacemakers; Technical				
ISO 5841-3 Technical	Con 2003-11	N	Corrigendum_1				

	1	T	Conical fittings with a 6 % (Luer) taper for syringes,		1	1		
			needles and certain other medical equipment; Part					
ISO 594-1	1986-06	N	1 : General requirements	Υ	N			AP3、4、23、34、40、41、43
30 394-1	1900-00	IN	Conical fittings with 6%_(Luer) taper for syringes,	1	IN			AF3, 4, 23, 34, 40, 41, 43
			needles and certain other medical equipment					
SO 594-2	1998-09	N	Part 2: Lock fittings	Υ	N			AD2 4 02 24 40 41 42
30 394-2	1990-09	IN	Reusable all-glass or metal-and-glass syringes for	I	IN			AP3、4、23、34、40、41、43
SO 595-1	1986-12	N	medical use; Part 1 : Dimensions					
30 393-1	1900-12	IN	Reusable all-glass or metal-and-glass syringes for					
			medical use; Part 2 : Design, performance					
SO 595-2	1987-12	N	requirements and tests					
50 393-Z	1907-12	IN	Hypodermic needles for single use; colour coding					
SO 6009	1992-12	N	for identification					
30 0009	1992-12	IN	ioi identinication					
			Hypodermic needles for single use - Colour coding					
SO 6009 Technical Corrig	2008-03	N						
30 0009 Technical Conig	2006-03	IN	for identification; Technical Corrigendum_1 Dentistry Number coding system for rotary instruments					
SO 6360-1	2004-04	N	Part_1: General characteristics					
			Dentistry Number coding system for rotary instruments					
			Part_1: General characteristics; Technical					
SO 6360-1 Technical Corrige	2007-09	N	Corrigendum_1					
			Dentistry Number coding system for rotary instruments					
SO 6360-2	2004-11	N	Part_2: Shapes					
			Dentistry Number coding system for rotary instruments Part 2: Shapes; Amendment 1					
SO 6360-2 AMD 1	2011-12	N	Part_2. Shapes, Amendment_1					
			Dentistry Number coding system for rotary instruments					
SO 6360-3	2005-11	N	Part 3: Specific characteristics of burs and cutters					
30 0300-3	2003-11	IN	Target opening arrangements of pare and called					
			Dentistry Number coding system for rotary instruments					
SO 6360-4	2004-06	N	Part_4: Specific characteristics of diamond instruments					
			Dentistry Number coding system for rotary instruments					
SO 6360-5	2007-12	N	Part_5: Specific characteristics of root-canal instruments					
			Dentistry Number anding quater for retary instruments					
SO 6360 6	2004-06	N	Dentistry Number coding system for rotary instruments Part 6: Specific characteristics of abrasive instruments					
SO 6360-6	2004-06	IN	Dentistry Number coding system for rotary instruments					
			Part_7: Specific characteristics of mandrels and special					
SO 6360-7	2006-02	N	instruments					
			Implants for surgery Ceramic materials Part_1:					
SO 6474-1	2010-02	N	Ceramic materials based on high purity alumina					
		† ''	2.2 materials susses on high party diamina		1	1		
		1	Implants for surgery; metal bone screws with					
		1	asymmetrical thread and spherical under-surface;					
SO 6475	1989-11	N	mechanical requirements and test methods	Υ	N			GL5、8
		1	Single-use containers for venous blood specimen	· · ·	1			, -
SO 6710	1995-08	N	collection					
SO 6872	2008-09	N	Dentistry Ceramic materials	Y	N			AP36
SO 6873	1998-03	N	Dental gypsum products					
SO 6874	2005-08	N	Dentistry - Polymer-based pit and fissure sealants	Р	N	JIS T 6524:2005	25.03.2005	275
00 00. 7	2000 00	1	y - - y	· ·	- ''	5.5 · 6024.2000	20.00.2000	
SO 6875	2011-07	N	Dentistry - Patient chair	P	N	JIS T 5602:2014	2014/3/1	160
ISO 6875	2011-07	N	Dentistry Patient chair	Р	N	JIS T 56 02: 2014	2014/3/1	160

ISO 6876	2001-08	N	Dental root canal sealing materials	Р	N	JIS T 6522:2005	25.03.2005	296
ISO 6877	2006-04	N	Dentistry Root-canal obturating points	Р	N	JIS T 6515:2011	29.07.2011	294
	2000 0 1	.,	Surgical instruments; non-cutting, articulated	· · · · · · · · · · · · · · · · · · ·		0.0 1 00 10.20 1	2010112011	
			instruments; general requirements and test					
ISO 7151	1988-12	N	methods					
	1000 12	.,	Surgical instruments; metallic materials; part_1:					
ISO 7153-1	1991-04	N	stainless steel	Υ	N			GL7、8
	100101		Surgical instruments Metallic materials Part_1:					927,0
ISO 7153-1 AMD 1	1999-03	N	Stainless steel; Amendment 1					
ISO 7176-1	1999-10	N	Wheelchairs - Part 1: Determination of static stability					
130 / 170-1	1999-10	IN	Wheelchairs Part_10: Determination of obstacle-					
ISO 7176-10	2008-11	N	climbing ability of electrically powered wheelchairs					
ISO 7176-11	1992-05	N	Wheelchairs; part 11: test dummies					
150 / 1/6-11	1992-05	IN	Wheelchairs; part_13: determination of coefficient of					
ISO 7176-13	1989-08	N	friction of test surfaces					
150 / 1/6-13	1909-00	IN	Wheelchairs - Part 14: Power and control systems for					
			electrically powered wheelchairs and scooters					
ISO 7176-14	2008-02	N	Requirements and test methods					
130 / 1/0-14	2000-02	IN	Wheelchairs Part_15: Requirements for information					
ISO 7176-15	1996-11	N	disclosure, documentation and labelling					
130 / 170-13	1990-11	IN	Wheelchairs Part_16: Resistance to ignition of					
ISO 7176-16	1997-05	N	upholstered parts Requirements and test methods					
100 / 1/0-10	1337-03	- 17	Wheelchairs - Part 19: Wheeled mobility devices for use					
ISO 7176-19	2008-07	N	as seats in motor vehicles					
100 7 170 10	2000 01	.,,	Wheelchairs Part_2: Determination of dynamic stability					
ISO 7176-2	2001-06	N	of electric wheelchairs					
100 1 11 0 2	200.00							
			Wheelchairs - Part 21: Requirements and test methods					
			for electromagnetic compatibility of electrically powered					
ISO 7176-21	2009-04	N	wheelchairs and scooters, and battery chargers					
ISO 7176-22	2000-05	N	Wheelchairs Part_22: Set-up procedures					
100 1 110 22	2000 00		Wheelchairs Part_23: Requirements and test methods					
ISO 7176-23	2002-07	N	for attendant-operated stair-climbing devices					
		1	Wheelchairs - Part 24: Requirements and test methods					
ISO 7176-24	2004-10	N	for user-operated stair-climbing devices					
ISO 7176-26	2007-04	N	Wheelchairs Part_26: Vocabulary					
100 1 110 20	2007 04	.,,	Wheelchairs Part_3: Determination of effectiveness of					
ISO 7176-3	2003-04	N	brakes					
100 1 110 0	2000 0 1		Wheelchairs Part_4: Energy consumption of electric					
			wheelchairs and scooters for determination of theoretical					
ISO 7176-4	2008-10	N	distance range					
			Wheelchairs Part_5: Determination of dimensions,					
ISO 7176-5	2008-06	N	mass and manoeuvring space					
					1			
			Wheelchairs Part_6: Determination of maximum speed,					
ISO 7176-6	2001-10	N	acceleration and deceleration of electric wheelchairs					
			Wheelchairs Part_7: Measurement of seating and					
ISO 7176-7	1998-05	N	wheel dimensions					
			Wheelchairs Part_8: Requirements and test methods					
ISO 7176-8	1998-07	N	for static, impact and fatigue strengths					
			Wheelchairs Part_9: Climatic tests for electric					
ISO 7176-9	2009-11	N	wheelchairs					
ISO 7193	1985-12	N	Wheelchairs: Maximum overall dimensions					

			N		1	
100 7407	0000 00		Neurosurgical implants Sterile, single-use	V	N	100
ISO 7197	2006-06	N	hydrocephalus shunts and components	Y	N	AP42
			Neurosurgical implants Sterile, single-use			
ISO 7197 Technical Corr	ia 2007 07	N	hydrocephalus shunts and components; Technical Corrigendum 1			
130 / 197 Technical Com	ig/2007-07	IN	Cardiovascular implants Tubular vascular			
ISO 7198	1998-08	N	prostheses			
100 7 190	1990-00	IN	Cardiovascular implants and artificial organs			
ISO 7199	2009-04	N	Blood-gas exchangers (oxygenators)	Υ	N	AP38
100 1100	2000 01	- 11	Cardiovascular implants and artificial organs -	•	.,	711 00
			Blood-gas exchangers (oxygenators)			
			Amendment_1: Clarifications for test			
ISO 7199 AMD 1	2012-02	N	methodologies, labelling, and sampling schedule			
			Implants for surgery Partial and total hip joint			
			prostheses Part_1: Classification and designation			
ISO 7206-1	2008-04	N	of dimensions			
			Implants for surgery Partial and total hip-joint			
l			prostheses Part_10: Determination of resistance			
ISO 7206-10	2003-12	N	to static load of modular femoral heads	Y	N	GL1
			Implants for surgery Partial and total hip joint			
100 7000 0	0044.04		prostheses Part_2: Articulating surfaces made of			
ISO 7206-2	2011-04	N	metallic, ceramic and plastics materials	Y	N	GL1
			Implants for surgery Partial and total hip joint prostheses Part_4: Determination of endurance			
			properties and performance of stemmed femoral			
ISO 7206-4	2010-06	N	components	Υ	N	GL1
100 1200 4	2010 00	1,	Implants for surgery; partial and total hip joint		11	GE1
			prostheses; part_6: determination of endurance			
			properties of head and neck region of stemmed			
ISO 7206-6	1992-03	N	femoral components	Υ	N	GL1
			Implants for surgery Components for partial and			
			total knee joint prostheses Part_1: Classification,			
ISO 7207-1	2007-02	N	definitions and designation of dimensions			
			Implants for surgery Components for partial and			
			total knee joint prostheses Part_2: Articulating			
100 7007 0	0044.07	N.	surfaces made of metal, ceramic and plastics	V	NI NI	01.0
ISO 7207-2	2011-07	N	materials	Y	N	GL2
ISO 7376	2009-08	N	Anaesthetic and respiratory equipment Laryngoscopes for tracheal intubation			
100 1010	2003-00	IN	Medical gas pipeline systems Part_1: Pipeline		+	
			systems for compressed medical gases and			
ISO 7396-1	2007-04	N	vacuum			
		1,				
			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			
ISO 7396-1 AMD 1	2010-01	N	connected to the pipeline through flexible hoses			

			Medical gas pipeline systems Part_1: Pipeline					
			systems for compressed medical gases and					
ISO 7396-1 AMD 2	2010-02	N	vacuum; Amendment 2					
			Medical gas pipeline systems Part_2: Anaesthetic					
ISO 7396-2	2007-04	N	gas scavenging disposal systems					
			Dentistry Evaluation of biocompatibility of medical	_			04.00.0040	0 //4/404) 01 /4
ISO 7405	2008-12	N	devices used in dentistry Copper-bearing contraceptive intrauterine devices	Р	N	JIS T 6001:2012	01.03.2012	See # 1 (161)on Sheet1
ISO 7439	2011-06	N	Requirements and tests					
ISO 7488	1991-06	N	Dental amalgamators					
ISO 7491	2000-09	N	Dental materials Determination of colour stability					
ISO 7492	1997-02	N	Dental explorers					
ISO 7493	2006-05	N	Dentistry Operator's stool					
100 / 100	2000 00	.,	Dentistry Dental units Part_1: General requirements					
ISO 7494-1	2011-08	N	and test methods	Р	N	JIS T 5701:2014	01.03.2014	160
ISO 7494-2	2003-03	N	Dentistry Dental units Part_2: Water and air supply Dental absorbent points					
ISO 7551	1996-12	N	Dental rotary instruments Diamond instruments					
			Part_1: Dimensions, requirements, marking and					
ISO 7711-1	1997-02	N	packaging					
			Dental rotary instruments Diamond instruments					
100 7744 4 AMD 4	2000 05		Part_1: Dimensions, requirements, marking and packaging; Amendment_1					
ISO 7711-1 AMD 1	2009-05	N	packaging, Amendment_1					
ISO 7711-2	2011-07	N	Dentistry Rotary diamond instruments Part_2: Discs					
			Dentistry Diamond rotary instruments Part_3: Grit					
ISO 7711-3	2004-11	N	sizes, designation and colour code					
			Instruments for surgery; Scalpels with detachable					
ISO 7740	1985-12	N	blades; Fitting dimensions					
100 7744	4000.00		Instruments for surgery; Scissors and shears;					
ISO 7741	1986-02	N	General requirements and test methods Dental handpieces Part_1: High-speed air turbine					
ISO 7785-1	1997-08	N	handpieces	Р	N	JIS T 5906:2001	25.05.2001	153
			Dental handpieces Part_2: Straight and geared angle					
ISO 7785-2	1995-08	N	handpieces	Р	N	JIS T5907:2011	29.07.2011	486 · 485 · 154
ISO 7786	0004.04	N	Dental rotary instruments Laboratory abrasive instruments					
150 7766	2001-04	IN	Dental rotary instruments; Cutters; Part 1 : Steel					
ISO 7787-1	1984-12	N	laboratory cutters					
			Dental rotary instruments Cutters Part_2: Carbide					
ISO 7787-2	2000-12	N	laboratory cutters					
100 7707 0	1001.10		Dental rotary instruments; cutters; part_3: carbide laboratory cutters for milling machines					
ISO 7787-3	1991-12	N	Dental rotary instruments Cutters Part_4: Miniature					
ISO 7787-4	2002-03	N	carbide laboratory cutters					
			·		1			64 · 65 · 67 · 69 · 72 · 507 · 508
								· 624 · 625 · 626 · 627 · 633 · 6
					1			34 · 637 · 671 · 679 · 685 · AP1
ISO 7864	1993-05	N	Sterile hypodermic needles for single use	Р	N	JIS T 3209:2011	29.07.2011	6
	1000 00	.,	, , , , , , , , , , , , , , , , , , ,		+ ''-	3.3 1 0200.2011	20.01.2011	<u> </u>
ISO 7885	2010-02	N	Dentistry - Sterile injection needles for single use	Р	N	JIS T 6130: 2013	01.09.2013	384

								64 · 72 · 628 · 630 · 632 · 638 ·
			Sterile hypodermic syringes for single use; part 1:					658 · 686 · 698 · AP37
SO 7886-1	1993-10	N	syringes for manual use	Υ	N			AP41· AP43
	1000			•				64 · 72 · 628 · 630 · 632 · 638 ·
			Sterile hypodermic syringes for single use - Part 1:					658 · 686 · 698 · AP37
ISO 7886-1 Technica	I Cort 1995-11	N	Syringes for manual use; Technical Corrigendum 1	Р	N	JIS T 3210:2011	29.07.2011	000 000 000 7.11 0.
100 7000 T Technica	1 0011 1333 11	14	by migos for manual abo, roomiscal comgonating.	<u>'</u>	14	010 1 3210.2011	25.07.2011	
			Sterile hypodermic syringes for single use - Part 2:					
ISO 7886-2	1996-05	N	Syringes for use with power-driven syringe pumps					
			Sterile hypodermic syringes for single use Part_3:					
ISO 7886-3	2005-03	N	Auto-disable syringes for fixed-dose immunization					
			Sterile hypodermic syringes for single use Part_4:					
ISO 7886-4	2006-10	N	Syringes with re-use prevention feature					
ISO 7944	1998-06	N	Optics and optical instruments Reference wavelengths					
150 7944	1998-06	IN	Optics and optical instruments Reference					
ISO 7944 Technical C	Corrig 2009-07	N	wavelengths; Technical Corrigendum 1					
100 70 11 1001111001	50111g 2000 07		Ophthalmic optics Spectacle frames Lists of					
ISO 7998	2005-10	N	equivalent terms and vocabulary					
			Mechanical contraceptives Reusable natural and					
			silicone rubber contraceptive diaphragms Requirements and tests					
ISO 8009	2004-10	N	Mechanical contraceptives - Reusable natural and					
			silicone rubber contraceptive diaphragms - Requirements					
ISO 8009 AMD 1	2012-02	N	and tests; Amendment_1					
			Small-bore connectors for liquids and gases in					
			healthcare applications Part_1: General					
ISO 80369-1	2010-12	N	requirements					
			Medical electrical equipment Part_2-12:					
100 00004 0 40	0044.04		Particular requirements for basic safety and					
ISO 80601-2-12	2011-04	N	essential performance of critical care ventilators Medical electrical equipment Part_2-12:					
			Particular requirements for basic safety and					
			essential performance of critical care ventilators;					
ISO 80601-2-12 Tech	nical 2011-10	N	Technical Corrigendum 1					
			Medical electrical equipment Part_2-13:					
			Particular requirements for basic safety and					
			essential performance of an anaesthetic					
ISO 80601-2-13	2011-08	N	workstation					
			Medical electrical equipment Part_2-55:					
ISO 80601-2-55	2011-12	N	Particular requirements for the basic safety and essential performance of respiratory gas monitors					
130 60601-2-33	2011-12	IN	Medical electrical equipment Part_2-56: Particular					
			requirements for basic safety and essential performance					
			of clinical thermometers for body temperature					
ISO 80601-2-56	2009-10	N	measurement					
			Medical electrical equipment Part_2-61:					
100 00004 0 04	2011 01	N.	Particular requirements for basic safety and					
ISO 80601-2-61	2011-04	N	essential performance of pulse oximeter equipment					1

		1			•	•		
			Non-invasive sphygmomanometers Part_1:					
			Requirements and test methods for non-automated					
ISO 81060-1	2007-12	N	measurement type					
			Non-invasive sphygmomanometers Part_2:					
ISO 81060-2	2009-05	N	Clinical validation of automated measurement type					
			Non-invasive sphygmomanometers Part_2:					
100 04000 0 T			Clinical validation of automated measurement type;					
ISO 81060-2 Techni	icai Co 2011-02	N	Technical Corrigendum_1 Respiratory tract humidifiers for medical use					
			Particular requirements for respiratory					
ISO 8185	2007-07	N	humidification systems	Υ	N	JIS T 7207:2005	25.03.2005	539· 540· 114
130 6163	2007-07	IN	Radiation protection; Clothing for protection against		IN	313 1 7207.2003	23.03.2003	559 540 114
			radioactive contamination; Design, selection, testing and					
ISO 8194	1987-06	N	use					
			Acoustics - Audiometric test methods - Part 1:					
ISO 8253-1	2010-11	N	Pure-tone air and bone conduction audiometry					
			Acoustics Audiometric test methods Part_2:					
			Sound field audiometry with pure-tone and narrow-					
ISO 8253-2	2009-12	N	band test signals					
			Acoustics Audiometric test methods Part_3:					
ISO 8253-3	2012-03	N	Speech audiometry					
			Dental equinoment Mercury and alloy mixers and					
ISO 8282	1994-10	N	dispensers					
			Orthopaedic instruments Drive connections					
100 0040 4	4000.05	N.	Part_1: Keys for use with screws with hexagon					
ISO 8319-1	1996-05	N	socket heads					
			Orthopaedic instruments; Drive connections; Part 2					
			: Screwdrivers for single slot head screws, screws					
ISO 8319-2	1986-10	N	with cruciate slot and cross-recessed head screws					
ISO 8325	2004-09	N	Dentistry - Test methods for rotary instruments					
100 0020	2004 00	14	Oxygen concentrators for medical use Safety					
ISO 8359	1996-12	N	requirements	Р	N	JIS T 7209:2007	25.01.2007	395
100 0000	1000 12		Injection containers and accessories Part_1:	<u> </u>	1,	010 1 7200.2007	20.01.2007	000
ISO 8362-1	2009-12	N	Injection containers and accessories1 art_1.					
100 0002 1	2000 12		Injection containers and accessories - Part 2:					
ISO 8362-2	2008-10	N	Closures for injection vials					
			Injection containers and accessories - Part 3:					
ISO 8362-3	2001-12	N	Aluminium caps for injection vials					
			Injection containers and accessories Part_4:					
ISO 8362-4	2011-09	N	Injection vials made of moulded glass					
			Injection containers and accessories Part_5:					
ISO 8362-5	2008-10	N	Freeze drying closures for injection vials					
			Injection containers and accessories Part_6:					
			Caps made of aluminium-plastics combinations for					
ISO 8362-6	2010-06	N	injection vials					
			Injection containers and accessories Part_7:					
			Injection caps made of aluminium-plastics					
ISO 8362-7	2006-04	N	combinations without overlapping plastics part					
100 0 400	1000.00		Optics and optical instruments; Ophthalmology;					
ISO 8429	1986-09	N	Graduated dial scale					

		Π	Infusion equipment for medical use Part_1:		1			1
ISO 8536-1	2011-09	N	Infusion equipment for medical use Part_1: Infusion glass bottles					
130 0330-1	2011-09	IN	Infusion grass bottles Infusion equipment for medical use Part_10:					
			Accessories for fluid lines for use with pressure					64 · 94 · 683 · 684 · 686 · 102 ·
ISO 8536-10	2004-10	N	infusion equipment	Р	N	JIS T 3211:2011	29.07.2011	AP13
150 0550-10	2004-10	11	Infusion equipment for medical use Part_11:	Г	in in	313 1 3211.2011	23.07.2011	Ai 13
			Infusion equipment for medical use_ rangers. Infusion filters for use with pressure infusion					
ISO 8536-11	2004-10	N	equipment	Р	N	JIS T 3219:2011	29.07.2011	102 · 683 · 684
130 6330-11	2004-10	IN	Infusion equipment for medical use Part_12:	г	IN	313 1 32 19.2011	29.07.2011	102 003 004
ISO 8536-12	2007-04	N	Check valves					
130 6330-12	2007-04	IN	Infusion equipment for medical use Part_2:					
ISO 8536-2	2010-03	N	Closures for infusion bottles					
100 0000-2	2010-03	IN	Infusion equipment for medical use - Part 3:					
ISO 8536-3	2009-06	N	Aluminium caps for infusion bottles					
100 0000 0	2003 00	14	Infusion equipment for medical use Part_4:					64 · 94 · 683 · 684 · 686 · 102 ·
ISO 8536-4	2010-10	N	Infusion sets for single use, gravity feed	Р	l N	JIS T 3211:2011	29.07.2011	AP13· 97
130 6536-4	2010-10	IN	9 19 7	<u> </u>	IN	JIS 1 3211.2011	29.07.2011	64 · 94 · 683 · 684 · 686 · 102 ·
			Infusion equipment for medical use Part_5:					
ISO 8536-5	2004-02	N	Burette infusion sets for single use, gravity feed	Р	N	JIS T 3211:2011	29.07.2011	AP13· 97
			Infusion equipment for medical use Part_6:					
ISO 8536-6	2009-11	N	Freeze drying closures for infusion bottles					
			Infusion equipment for medical use Part_7: Caps					
			made of aluminium-plastics combinations for					
ISO 8536-7	2009-01	N	infusion bottles					
			Infusion equipment for medical use Part_8:					64 · 94 · 683 · 684 · 686 · 102 ·
			Infusion equipment for use with pressure infusion	_				
ISO 8536-8	2004-08	N	apparatus	Р	N	JIS T 3211:2011	29.07.2011	AP13 · 97
			Inferior and instantion and include Deat O. Floid					64 · 94 · 683 · 684 · 686 · 102 ·
			Infusion equipment for medical use Part_9: Fluid					
ISO 8536-9	2004-10	N	lines for use with pressure infusion equipment	Р	N	JIS T 3211:2011	29.07.2011	AP13· 97
			Sterile single-use syringes, with or without needle,					
ISO 8537	2007-10	N	for insulin	Р	N	JIS T 3253:2012	01.07.2012	AP29
			Prosthetics and orthotics; limb deficiencies; part_1:					
			method of describing limb deficiencies present at					
ISO 8548-1	1989-08	N	birth					
100 0540 0	4000.07	N	Prosthetics and orthotics; limb deficiencies; part_2:					
ISO 8548-2	1993-07	N	method of describing lower limb amputation stumps					
			Prosthetics and orthotics; limb deficiencies; part_3: method of describing upper limb amputation					
ISO 8548-3	1993-07	N	stumps					
130 6546-3	1993-07	IN	Prosthetics and orthotics Limb deficiencies					
			Part 4: Description of causal conditions leading to					
ISO 8548-4	1998-07	N	amputation					
100 00 10 1	1000 07	.,	Prosthetics and orthotics - Limb deficiencies -					
			Part 5: Description of the clinical condition of the		1			
ISO 8548-5	2003-07	N	person who has had an amputation		1			
			i i					
			Prosthetics and orthotics; vocabulary; part_1: general					
ISO 8549-1	1989-07	N	terms for external limb protheses and external orthoses					
			Prosthetics and orthotics; vocabulary; part_2: terms					
100 0540 0	4000.07		relating to external limb prostheses and wearers of these prostheses		1			
ISO 8549-2	1989-07	N	prostreses					

			Prosthetics and orthotics; vocabulary; part_3: terms					
ISO 8549-3	1989-07	N	relating to external orthoses					
			Prosthetics and orthotics Functional deficiencies					
			Description of the person to be treated with an orthosis, clinical objectives of treatment, and functional					
ISO 8551	2003-08	N	requirements of the orthosis					
130 0331	2003-00	IN	Ophthalmic optics Visual acuity testing					
ISO 8596	2009-07	N	Standard optotype and its presentation					
130 0390	2009-07	IN	Standard optotype and its presentation					MHLW Ministerial Ordinance No.
ISO 8598	1996-08	N	Optics and optical instruments - Focimeters	Р	Р	JIS B7183:1995	01.12.1995	349,2001
			Optics and optical instruments Focimeters;					
ISO 8598 Technical Corri	1998-05	N	Technical corrigendum_1					
			Optics and photonics Medical endoscopes and					55 · 56 · 57 · 58 · 62 · 63
			endotherapy devices Part_1: General					501 · 503 · 504 · 521 · 620 · AP
ISO 8600-1	2005-05	N	requirements	Р	N	JIS T 1553:2005	25.03.2005	39,40
130 0000-1	2003-03	IN	roquiromonio	г	IN	JIS 1 1555.2005	23.03.2003	39,40
1			Optics and optical instruments Medical					
			endoscopes and endoscopic accessories - Part 2:					
ISO 8600-2	2002-08	N	Particular requirements for rigid bronchoscopes					
130 8000-2	2002-00	IN	Optics and optical instruments - Medical					
			endoscopes and endoscopic accessories Part_3:					55 · 56 · 57 · 58 · 62 · 63
			Determination of field of view and direction of view					501 · 503 · 504 · 521 · 620 · AP
ISO 8600-3	1997-07	N	of endoscopes with optics	Р	N	JIS T 1553:2005	25.03.2005	39,40
130 0000-3	1331-01	IN	or oridocoopee man opine	r	IN .	313 1 1333.2003	23.03.2003	39,40
			Ontice and enticel instruments Medical					
			Optics and optical instruments Medical endoscopes and endoscopic accessories Part_3:					
			Determination of field of view and direction of view					
ISO 8600-3 AMD 1	2003-12	N	of endoscopes with optics: Amendment 1					
130 8000-3 AMD 1	2003-12	IN	Optics and optical instruments Medical					
			endoscopes and certain accessories - Part 4:					
			Determination of maximum width of insertion					55 · 56 · 57 · 58 · 62 · 63
ISO 8600-4	1997-07	N	portion	Р	N	JIS T 1553:2005	25.03.2005	501 · 503 · 504 · 521 · 620
150 0000-4	1997-07	IN.	portion	г	IN .	313 1 1333.2003	23.03.2003	301 303 304 321 020
			Optics and photonics Medical endoscopes and					
			endotherapy devices Part_5: Determination of					
ISO 8600-5	2005-03	N	optical resolution of rigid endoscopes with optics	Υ	N			AP39、40
130 0000-3	2003-03	IN	Optics and photonics - Medical endoscopes and	<u> </u>	IN .			AF 33, 40
ISO 8600-6	2005-03	N	endotherapy devices - Part 6: Vocabulary					
ISO 8612	2009-10	N	Ophthalmic instruments - Tonometers	Р	N	JIS T 7312:2005	25.03.2005	38-610
.00 00.2	2000 .0		Implants for surgery; fixation devices for use in the		.,	0.0 1 1012.2000	20.00.2000	00 0.0
ISO 8615	1991-11	N	ends of the femur in adults					
			Ophthalmic optics Spectacle frames Measuring					
ISO 8624	2011-02	N	system and terminology					
			Cardiovascular implants and extracorporeal					
			systems - Haemodialysers, haemodiafilters,					AP4
ISO 8637	2010-07	N	haemofilters and haemoconcentrators	Υ	N			AP17
			Cardiovascular implants and extracorporeal					
			systems Extracorporeal blood circuit for					
ISO 8638	2010-07	N	haemodialysers, haemodiafilters and haemofilters	Р	N	JIS T 3248: 2012	2012/10/1	97 · 793
ISO 8669-1	1988-07	N	Urine collection bags; part 1: vocabulary					

		_						1
100 0000 0	1000 10		Urine collection bags Part_2: Requirements and test methods					
ISO 8669-2	1996-12	N	Ostomy collection bags; part_1: vocabulary					
ISO 8670-1	1988-07	N	Ostomy collection bags, part_1. vocabulary Ostomy collection bags Part_2: Requirements and test					
ISO 8670-2	1996-12	N	methods					
100 0070 2	1000 12	- "	Ostomy collection bags Part_3: Determination of odour					
ISO 8670-3	2000-03	N	transmission of colostomy and ileostomy bags					
			Implants for surgery; staples with parallel legs for					
ISO 8827	1988-10	N	orthopaedic use; general requirements					
			Implants for surgery; guidance on care and					
ISO 8828	1988-10	N	handling of orthopaedic implants					
			Inhalational anaesthesia systems Part_7:					
			Anaesthetic systems for use in areas with limited					
			logistical supplies of electricity and anaesthetic					
ISO 8835-7	2011-11	N	gases					
ISO 8836	2007-09	N	Suction catheters for use in the respiratory tract	Р	N	JIS T 3251:2011	29.07.2011	83 · 644
			Elastomeric parts for parenterals and for devices					
100 0074 4	0000 40		for pharmaceutical use Part_1: Extractables in					
ISO 8871-1	2003-10	N	aqueous autoclavates Elastomeric parts for parenterals and for devices					
			for pharmaceutical use - Part 2: Identification and					
ISO 8871-2	2003-10	N	characterization					
100 007 1 2	2000 10	- ''	GHARACICHIZATION					
			Elastomeric parts for parenterals and for devices					
			for pharmaceutical use Part_2: Identification and					
ISO 8871-2 AMD 1	2005-07	N	characterization; Amendment_1					
			Elastomeric parts for parenterals and for devices					
			for pharmaceutical use Part_3: Determination of					
ISO 8871-3	2003-08	N	released-particle count					
			Elastomeric parts for parenterals and for devices					
			for pharmaceutical use Part_4: Biological					
ISO 8871-4	2006-06	N	requirements and test methods					
			Elastomeric parts for parenterals and for devices					
ISO 8871-5	2005-08	N	for pharmaceutical use Part_5: Functional requirements and testing					
130 007 1-3	2005-06	IN	Aluminium caps for transfusion, infusion and					
			injection bottles General requirements and test					
ISO 8872	2003-03	N	methods					
	2000 00	.,	Ophthalmic optics Uncut finished spectacle					
			lenses Part_1: Specifications for single-vision					
ISO 8980-1	2004-02	N	and multifocal lenses					
			Ophthalmic optics Uncut finished spectacle					
			lenses Part_1: Specifications for single-vision					
ISO 8980-1 Technical Cor	rı 2006-08	N	and multifocal lenses; Technical Corrigendum_1					
			Ophthalmic optics Uncut finished spectacle					
		1	lenses Part_2: Specifications for progressive					
ISO 8980-2	2004-02	N	power lenses					
			Ophthalmic optics Uncut finished spectacle					
ICO 0000 2 Tooks:! O	2000 00	l N	lenses Part_2: Specifications for progressive					
ISO 8980-2 Technical Co.	II ∠UUb-U8	N	power lenses; Technical Corrigendum_1					

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			Ophthalmic optics Uncut finished spectacle					
			lenses Part_3: Transmittance specifications and					
ISO 8980-3	2003-10	N	test methods					
			Ophthalmic optics Uncut finished spectacle					
ISO 8980-4	2006-08	N	lenses Part_4: Specifications and test methods for anti-reflective coatings					
150 8980-4	2006-08	IN	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					
100 0000			Dentistry Hose connectors for air driven dental					
ISO 9168	2009-07	N	handpieces					
			Terminal units for medical gas pipeline systems					
			Part_1: Terminal units for use with compressed					
ISO 9170-1	2008-07	N	medical gases and vacuum					
			Terminal units for medical gas pipeline systems					
100 0470 0	0000 07	N.	Part_2: Terminal units for anaesthetic gas					
ISO 9170-2	2008-07	N	scavenging systems Dentistry Extraction forceps Part_1: General					
ISO 9173-1	2006-06	N	requirements and test methods					
ISO 9173-2	2010-05	N	Dentistry Extraction forceps Part_2: Designation					
			Injection equipment for medical use Part_1:					
ISO 9187-1	2010-10	N	Ampoules for injectables					
			Injection equipment for medical use Part_2: One-					
ISO 9187-2	2010-10	N	point-cut (OPC) ampoules					
			Implants for surgery; metal bone screws with					
ISO 9268	1988-12	N	conical under-surface of head; dimensions					
			Implants for surgery; metal bone plates; holes and					
100 0000	1000.10	NI NI	slots corresponding to screws with conical under-					
ISO 9269	1988-12	N	surface			UO T 0447 0044	2011/7/29	190
ISO 9333	2006-07	N	Dentistry - Brazing materials	Р	N	JIS T 6117:2011 JIS T 6111:2011	2011/7/29	195
100 3333	2000-07	11	Dentistry Brazing materials	'	11	310 1 0111.2011	2011/1/25	133
			Optics and optical instruments - Test lenses for					
			calibration of focimeters Part_1: Test lenses for					
ISO 9342-1	2005-05	N	focimeters used for measuring spectacle lenses					
			Optics and optical instruments Test lenses for					
			calibration of focimeters Part_2: Test lenses for					
ISO 9342-2	2005-11	N	focimeters used for measuring contact lenses					
			Anaesthetic and respiratory equipment - Heat and					
			moisture exchangers (HMEs) for humidifying					
			respired gases in humans Part_1: HMEs for use					
ISO 9360-1	2000-03	N	with minimum tidal volumes of 250 ml					
			Anaesthetic and respiratory equipment - Heat and					
			moisture exchangers (HMEs) for humidifying					
			respired gases in humans Part_2: HMEs for use					
			with tracheostomized patients having minimum					
ISO 9360-2	2001-04	N	tidal volumes of 250_ml					
			Power-operated lifting platforms for persons with impaired mobility Rules for safety, dimensions and functional					
ISO 9386-1	2000-11	N	operation Part_1: Vertical lifting platforms					
.55 5555 1	2000-11	1 14	,		1			

				1			
		Power-operated lifting platforms for persons with impaired					
		mobility Rules for safety, dimensions and functional					
		operation Part_2: Powered stairlifts for seated, standing					
2000-11	N	and wheelchair users moving in an inclined plane					
		Ophthalmic optics Contact lenses and contact					
		lens care products Determination of					
1998-08	N	biocompatibility by ocular study with rabbit eyes					
1993-10	N						
		1 37,					
1000 10							
1993-10	N						
1000 12	N		V	N			GL5、8
1990-12	IN		t	IN			GL5, 8
1991-09	N	S S	V	N			AP16
1551 65	11			- 11			AT TO
2001-06	N						
2007-06	N	Dentistry Operating lights					
	N	Dental equipment; graphical symbols					
1999-12	N	Metal-ceramic dental restorative systems	Р	N	JIS T 6516:2005	2005/3/25	222
2005 10		, in the second	D	N	IIC T 6516:2005	25.03.2005	222
2003-10	IN	, , , –		111	313 1 03 10.2003	25.05.2005	222
2012-02	N	systems					
		Neurosurgical implants - Self-closing intracranial					
2002-09	N	aneurysm clips	Υ	N			AP27·AP28
		Orthopaedic drilling instruments; part_1: drill bits,					
1991-03	N	taps and countersink cutters					
2009-12	N	Ophthalmic instruments Trial case lenses					
1998-11	N	Dental hand instruments Reusable mirrors and handles					
		Dental hand instruments Reusable mirrors and handles;					
rrigend 2000-06	N	Technical Corrigendum_1					
		Dentistry Water-based cements Part_1:					253 · 254 · 258 · 271 · 272 · 274
2007-10	N	Powder/liquid acid-base cements	Р	N	JIS T 6609-1:2005	25.03.2005	· 276
		Dentistry Water-based cements Part_2: Resin-					
2010-04	N	modified cements	Р	N	JIS T 6609-2: 2014	2014/3/1	262 · 273 · 277 · 281
		,					
1993-07	N						
1002.07	N.						
1999-12	N						
2011-07	N	Classification and terminology					
	1998-08 1993-10 1993-10 1993-10 1990-12 1991-09 2001-06 2007-06 1993-02 1999-12 2005-10 2012-02 2002-09 1991-03 2009-12 1998-11 rrigen 2000-06 2007-10 2010-04 1993-07 1993-07 1993-07 1999-12	1998-08 N 1993-10 N 1993-10 N 1993-10 N 1990-12 N 1991-09 N 2001-06 N 2007-06 N 1993-02 N 1993-02 N 2002-09 N 1991-03 N 2002-09 N 1991-03 N 2009-12 N 1998-11 N rrigen(2000-06 N 2007-10 N 2010-04 N 1993-07 N 1993-07 N 1993-07 N 1993-07 N 1993-07 N	operation Part. 2: Powered stairlifts for seated, standing and wheelchair users moving in an inclined plane Ophthalmic optics Contact lenses and contact lens care products Determination of biocompatibility by ocular study with rabbit eyes Implants for surgery; non-destructive testing; liquid penetrant inspection of metallic surgical implants Implants for surgery; non-destructive testing; radiographic examination of cast metallic surgical implants Implants for surgery; non-destructive testing; radiographic examination of cast metallic surgical implants Implants for surgery; determination of bending strength and stiffness of bone plates Stainless steel needle tubing for manufacture of medical devices Stainless steel needle tubing for the manufacture of medical devices; Amendment_1 2001-06 N Dentistry Operating lights 1993-02 N Dental equipment; graphical symbols 1999-12 N Metal-ceramic dental restorative systems 2005-10 N Metal-ceramic dental restorative systems; Amendment_1 Dentistry Compatibility testing Part_1: Metal-ceramic systems 2002-09 N Autorosurgical implants Self-closing intracranial aneurysm clips Orthopaedic drilling instruments; part_1: drill bits, taps and countersink cutters 2009-12 N Dental hand instruments Reusable mirrors and handles Dental hand instruments_	mobility_ Rules for safety, dimensions and functional operation_ Part_2. 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