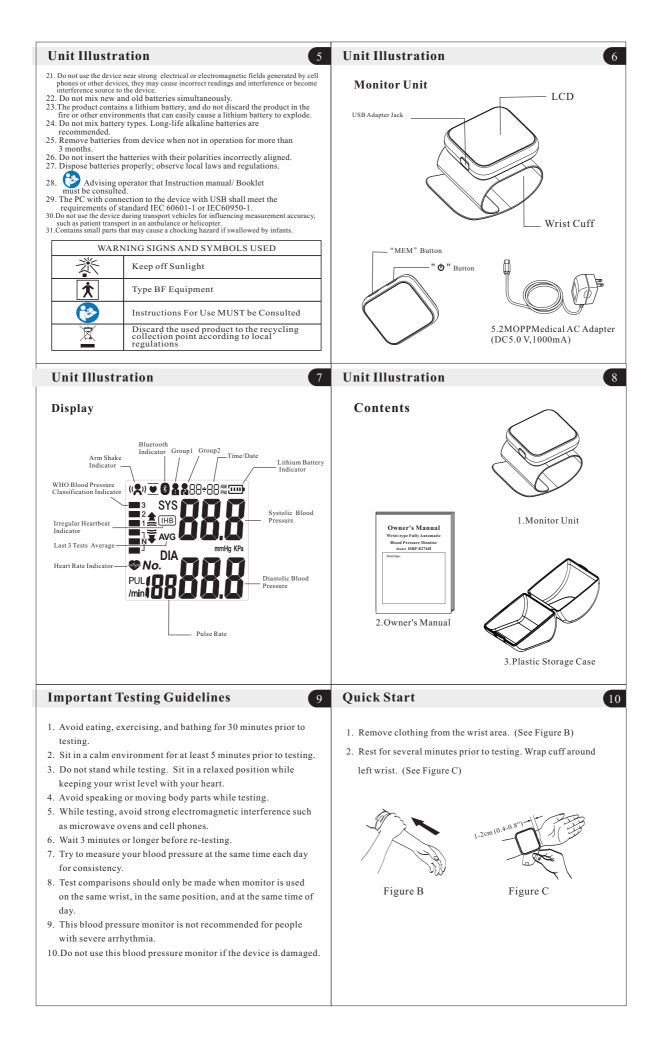
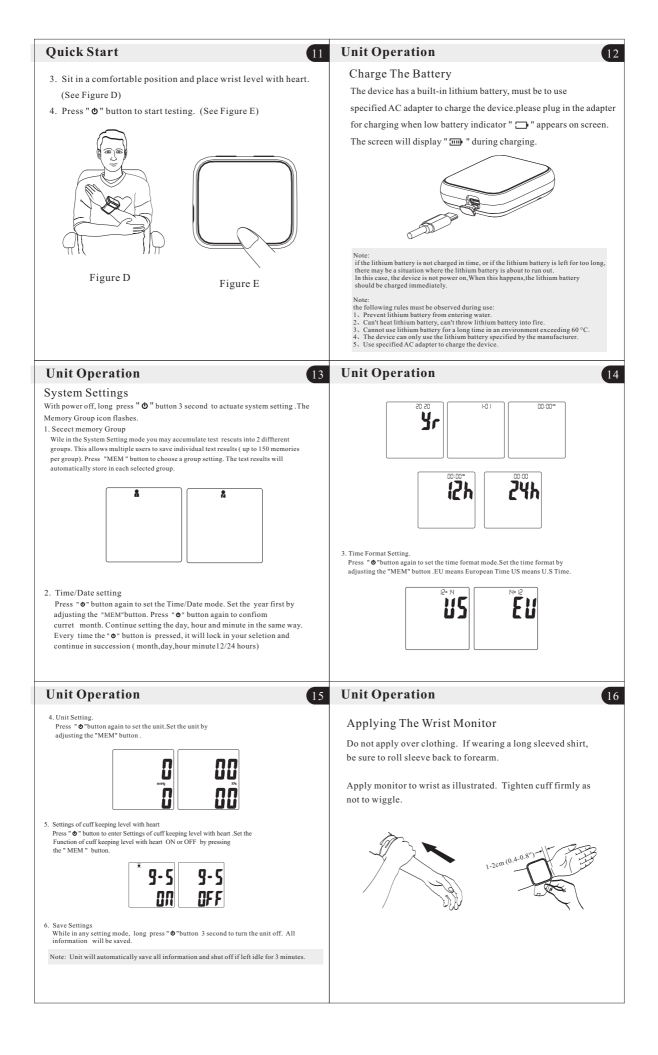
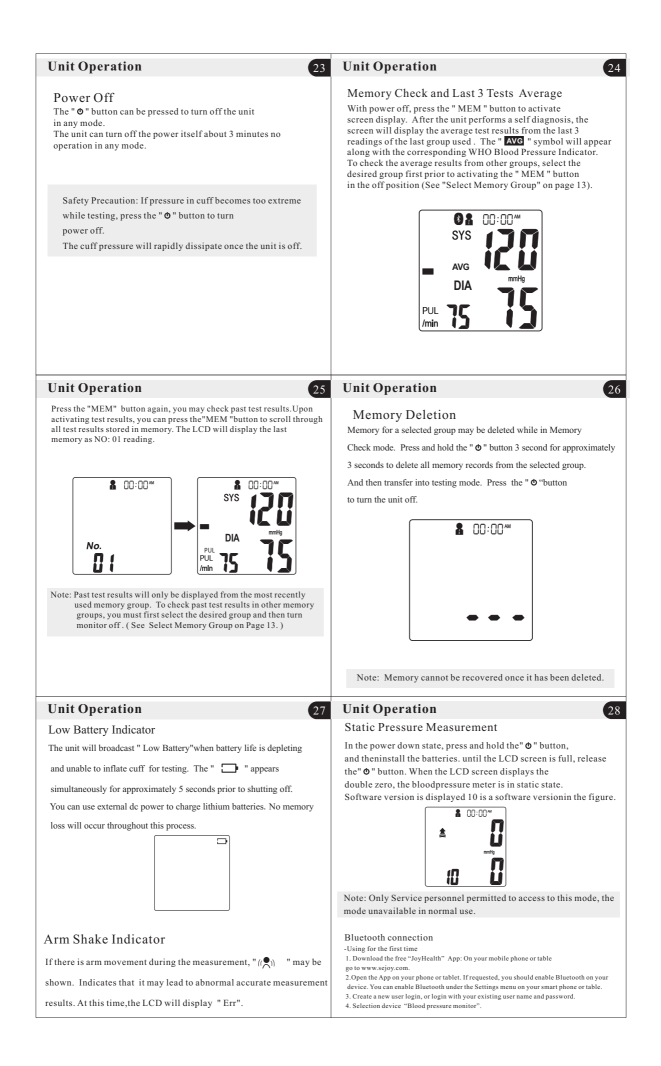
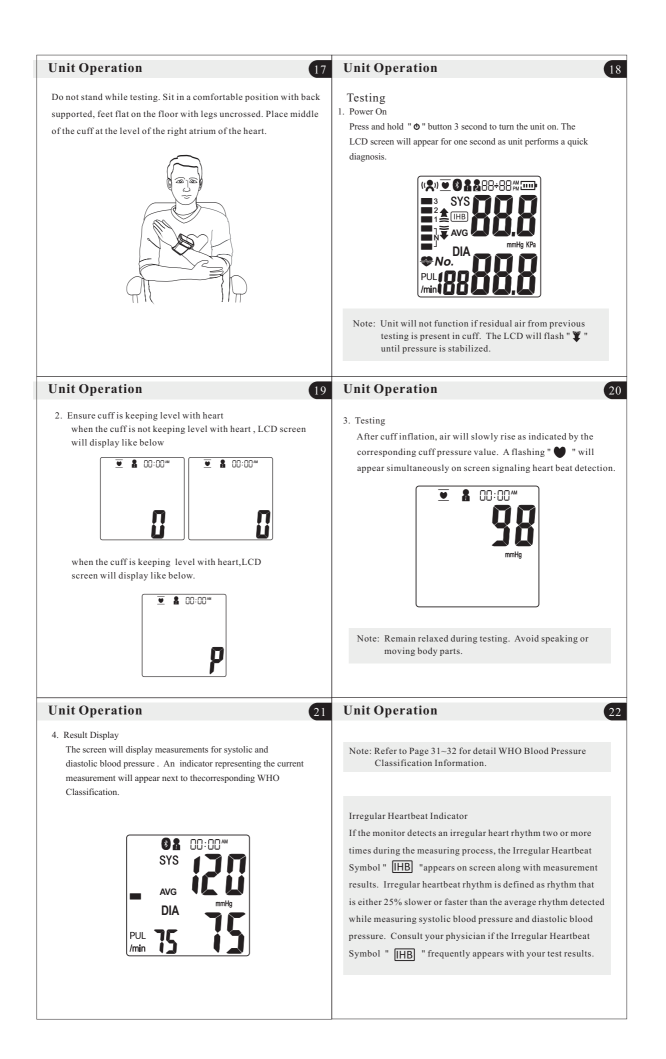


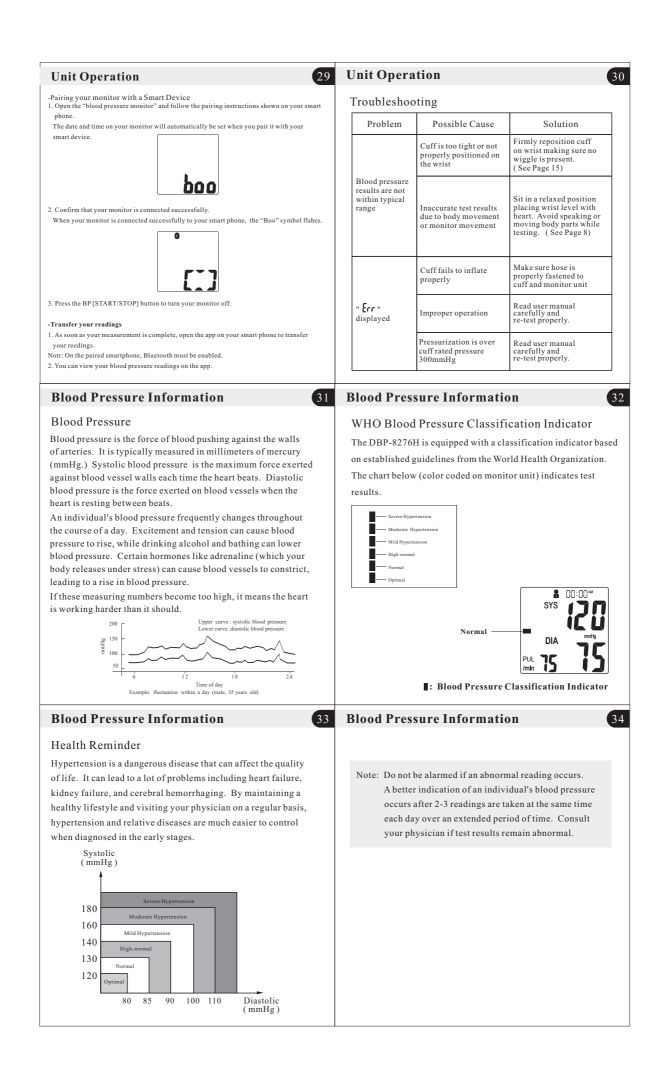
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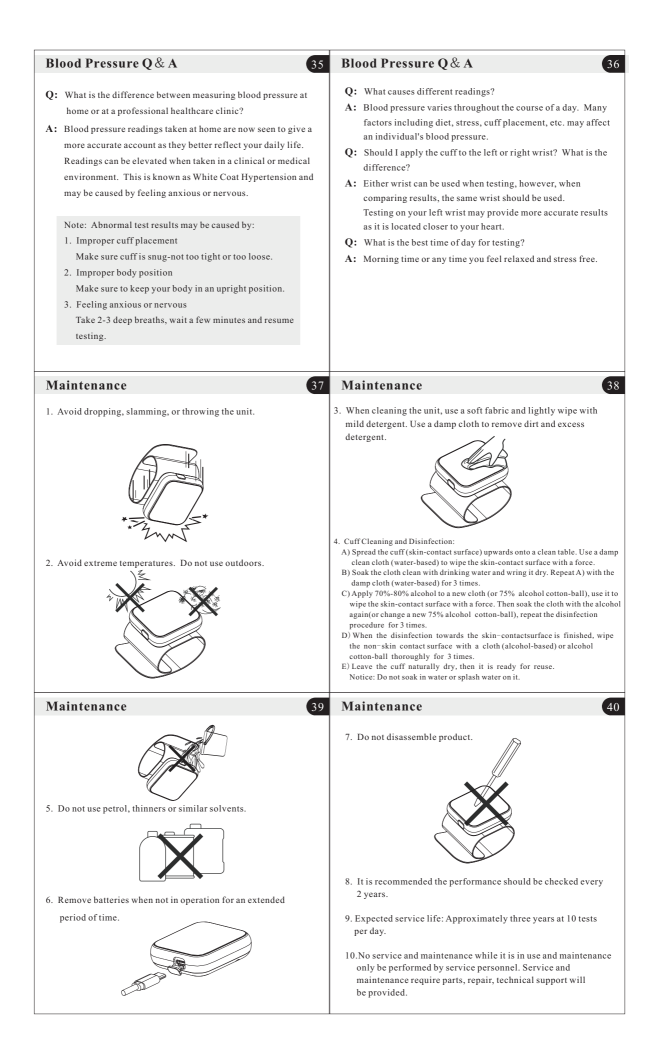












	5		4	1 Sj	pecifications				
Product Description	Wrist-Type Fully Automatic] [Function	Buzzer			
*		Digital Blood Pressure Monitor		41 F	Damas Sa	Backlight			
Model Display	DBP-8276H LCD Digital Display Size:34mm×34.6mm (1.34" x 1.36")		4 I F	Power Source	lithium battery (3.7v 180mAh)				
Measurement Method		Oscillometric Method			Battery run time	500 measurements			
~	Systolic Pres	Systolic Pressure 60mmHg~260mmHg			Charge time Unit Weight	1 hours Approx. 122g (4.300z.) (Excluding Battery)			
Measurement Range	Diastolic Pres	ssure 30	mmHg~20ommHg	7 I F	Unit Dimensions	Approx. 62mm×55.2mm×19mm(L x W x H) (2.44 " x 2.17"x 0.75")			
	Pressure	0n	nmHg~299mmHg	7 I F	Cuff Circumference	Fits wrist circumference 13.5-21.5 cm(5			
	Pressure	±3	mmHg			Temperature	$10^{\circ}C \sim 40^{\circ}C (50^{\circ}F \sim 104^{\circ}F)$		
	Pulse	30	~ 180 Beats/Minute] ,	Operating Environment	Humidity	15%~93%RH		
	Pulse	±5	±5%		-	Pressure	800hPa~1060hPa		
Pressurization	Automatic Pr	essurization		_		Temperature	-25°C~55°C (-13°F~131°F)		
Memory	2x60 Memories in Tow Groups with Date and Time]	Stamon E	Humidity	≤93% RH		
	Irregular Hea	Irregular Heartbeat Detection			Storage Environment Transport Environment	Temperature			
	WHO Classification Indicator			1 :					
Function	Last 3 Result	s Average		1 L		Humidity	≤93% RH		
	Low Battery	Detection		11					
	Automatic Po	Automatic Power-Off							
Specifications	5		4	3 Sj	pecifications				
	Modulation Type	GFSK		Sp	ecifications are subj	ect to chang	ze without notice.		
Bluetooth	Version	5.0.1 BT Signal mode			is Blood Pressure M	onitor com	plies with the European regula		
	Operation		(2400 [~] 2483. 5MHz)	and		CE 0197".	This blood pressure monitor al		
	frequency Antenna gain			(in	cluded but not limite	ed):	anuarus		
	-	0.5 dBi		- EN			uipment part 1: General		
	Bandwidth	2.0 MH	z	EN	uirements for safety IC standard:				
Ingress Protection Rating					EN 60601-1-2 Medical Electrical Equipment Part 1-2: General Requirements For Basic Safety And Essential Performance Collateral Standard: Electromagnetic Disturbances -				
Classification	Internal Powered Equipment Type BF								
Battery Shelf life	60 months				Requirements And Tests.				
Battery Storage Temperature	-25℃~55℃ ((-13°F~13	l°F)	IEC		al electrica	al equipment – Part 2-30: Partic		
Battery Storage	-25°C~55°C ((-13°F~13	"F)	IEC req aut EN req sys ISC	C80601-2-30, Medic puirements for the ba tomated non-invasiv 1 1060-3 Non-invasiv puirements forelectro stems.	eal electrica sic safety a e sphygmory we sphygmory omechanica sive sphygn	nd essential performance of manometers. omanometers - Supplementary Il blood pressure measuring momanometers - part 2: clinica		
Battery Storage	-25°C~55°C ((-13°F~13		S E	C80601-2-30, Medic uirements for the ba iomated non-invasiv 1060-3 Non-invasiv juirements forelectro stems. 0 81060-2, non-inva lidation of automated lectromagnetic	eal electrica sic safety a e sphygmore omechanica sive sphygm d measurem	nd essential performance of manometers. Supplementary il blood pressure measuring momanometers - part 2: clinica nent type.		
Battery Storage Temperature Warranty The Blood Pressure M late of purchase. If th properly due to defect will repair or replace	Monitor is gua he Blood Pres tive compone it freely. The	aranteed f isure Mor ents or po e warrant	For 2-year from the hitor does not function or workmanship, we	-5 E	C80601-2-30, Medic uirements for the ba tomated non-invasiv 1060-3 Non-invasiv uirements forelectro stems. D 81060-2, non-inva lidation of automated lectromagnetic he device satisfies the EM 0601-1-2. The requiremen ble below. The device is an recautionary measures with istructions for use. Portabl he device. Use of the unit is a device negatively and a hould not be used directly is ble 1	eal electrica sic safety a e sphygmo omechanica sive sphygm d measurem Compati Crequirement ts are satisfied n electrical me h regard to EM le and mobile I n conjunction ler the electro adjacent to or b	nd essential performance of manometers. sumanometers - Supplementary il blood pressure measuring momanometers - part 2: clinica ent type. (bility Information) s of the international standard IEC under the conditions described in the dical product and is subject to special IC which must be published in the HF communications equipment can affee with non-approved accessories can affe magnetic compatibility. The device between other electrical equipment.		
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Battery Storage Temperature Warranty The Blood Pressure M late of purchase. If th properly due to defect vill repair or replace lamages to your Bloo	Monitor is gua he Blood Pres stive compone it freely. The od Pressue Mo	aranteed f sure Mor ents or po e warrant onitor du	For 2-year from the hitor does not function or workmanship, we y does not cover	-5 E	C80601-2-30, Medic uirements for the ba tomated non-invasiv 1060-3 Non-invasiv stems. D 81060-2, non-inva lidation of automated lectromagnetic he device satisfies the EM 0601-1-2. The requiremen ble below. The device is ar recautionary measures wit nstructions for use. Portabl he device. Use of the unit i to edvice negatively and a hould not be used directly is ble 1 Guidance and declaration of The device is intended for us The customer or the user of the Emissions test	al electrica sic safety a e sphygmot omechanica sive sphygmo omechanica sive sphygmo d measurem Compati C requirement ts are satisfied n celetrical me h regard to EM le and mobile I n conjunction Iter the electro adjacent to or t f manufacturer- f manufacturer- se in the electron he device should Group I, ClassB	nd essential performance of manometers. manometers - Supplementary il blood pressure measuring momanometers - part 2: clinica tent type. Example 1 Supplementary is the supplementary of the suppl		
Battery Storage Temperature Warranty The Blood Pressure M late of purchase. If th properly due to defect will repair or replace lamages to your Bloo	Monitor is gua he Blood Pres stive compone it freely. The od Pressue Mo	aranteed f sure Mor ents or po e warrant onitor du	For 2-year from the hitor does not function or workmanship, we y does not cover	-5 E	C80601-2-30, Medic uirements for the ba tomated non-invasiv (1060-3 Non-invasiv uirements forelectro stems. D 81060-2, non-inva lidation of automated lectromagnetic he device satisfies the EM 0601-1-2. The requiremen blo below. The device is a recautionary measures with structions for use. Portable he device. Use of the unit i the device. Use of the unit is dould not be used directly and a hould not be used directly ble 1 Guidance and declaration of The device is intended for us The customer or the user of fl Emissions test Radiated emission CISPR 11	al electrica sic safety a e sphygmot omechanica sive sphygmo omechanica sive sphygmo d measurem Compati C requirement ts are satisfied n celetrical me h regard to EM le and mobile I n conjunction Iter the electro adjacent to or t f manufacturer- f manufacturer- se in the electron he device should Group I, ClassB	nd essential performance of manometers. manometers - Supplementary il blood pressure measuring momanometers - part 2: clinicate tent type. Example 1 So the international standard IEC under the conditions described in the dical product and is subject to special IC which must be published in the HF communications equipment can affer with non-approved accessories can affer magnetic compatibility. The device between other electrical equipment. Electromagnetic environment electromagnetic environment electr		

RC 60601 C RC 60601 C Start Level 1 S kV S kV S kV S kV S kV S kV S kV	evel $\pm 8 \text{ kV}$ contact $\pm 2 \text{ kV}, \pm 4 \text{ kV}, \text{ contact}$ $\pm 8 \text{ kV}, \pm 1 \text{ kV}, \text{ contact}$	c environment s that it is used in Electromagneti -galdance Floors should be or ceramic tile. I. overed with sy- overed with sy- overed with sy- not should be overed with sy- that should be in the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the system of the s	such an enviror c environment wood, concrete f floors are ithetic material,		ent. IMMUNITY test	IEC (0(01	Compliance level 3V/m or 10 V/m 80MHz-2.7 Ghz 80%AM at	
set level it ontact : ontact : ontact : ontact : ontact : 15 kV vair : 12 kV for : www.supply : 12 kV for : 12 kV for : 12 kV for : int kV for : put/output : 12 kV for : 04 differential : 25% UT : 05% UT : : : : : : : : : : : : : : : : : : : : : : : :	evel ± 8 kV contact ± 2 kV, ± 4 kV, e ± 2 kV, ± 4 kV, e ± 15 kV air N/A N/A	-guidance Floors should be or ceramic tile. It covered with syn the relative humi	wood, concrete f floors are thetic material,		Radiated RF EM fields	1 3V/m or 10 V/m 80MHz-2.7 Gbz 80%AM at	3V/m or 10 V/m 80MHz-2.7 Ghz 80%AM at	Portable and mobile RF communications equipment should be used no closer to an part of the device, including cables, than the recommended separation distance calculated from the equation applicable the frequency of the transmitter. Recommended separation distance of the second second second stance of the second seco
11 sk V air 21 sk V air 21 kV for www.supply. 21 kV for put/output. 10 kV for 10	×15 kV air s	he relative humi	dity should be	_	fields	80MHz-2.7 Ghz 80%AM at	80MHz-2.7 Ghz 80%AM at	calculated from the equation applicable the frequency of the transmitter. Recommended seperation distance 80 M to 800 MHz 800 MHz to 2.7 Gbz where F
nes nes 1 kV fifte iffer 2 kV sommon sommon sommon 5% UT 5% UT 5% UT 50% dip in 71 for 0.5 ycle 0% UT 50% dip in 71 for 25 ycle 5% UT 71 for 5 secretary	N/A			_		lkHz		the maximum output power rating of the transmitter in watts (W) according to the
20mmon 100de 5% UT -95% dip in TJ for 0.5 ycle 0% UT 0% dip in TJ for 5 ycle 0% UT 0% dip in TJ for 25 ycle 5% UT -95% dip in TJ for 5 scretary							1kHz	transmitter manufacturer and d is the recommended separation distance in metres (m). Field strengths from fixed R transmitters, as determined by an electromagnetic site survey, a should be less than the compliance level in each
ycle 0% UT 1% for 5 ycle 0% UT 80% dip in 17 for 25 ycle 5% UT -95% dip in T) for 5 scretary	N/A							frequency range. Interference may occur the vicinity of equipment marked with th following symbol: Portable and mobile RF communication equipment should be used no closer to a part of the device, including cables, than
					Conducted disturbances Induced by RF fields IEC 61000-4-6	3 V in 0.15 MHz- 80 MHz 6 V in ISM and/or amateur radio bands between 0.15 MHz and 80 MHz 80 % AM at 1kHz	3 V in 0.15 MHz- 80 MHz 6 V in ISM and/or amateur radio bands between 0.15 MHz and 80 MHz 80 % AM at 1kHz	recommended separation distance calculated from the equation applicable Recommended separation distance 80 M to 800 MHz 800 MHz to 2.7 Ghe where the maximum output power rating of the transm-itter manufacturer and di si the recommended separation distance in metres ((1), Field strengths from God R electromagnetic site survey, a should be han the compliance level in each freque vicinity of equipment marked with the following symbols 1.3
	30 A/m; 50Hz	Power frequency should be at leve typical location rcial or hospital	ils charactertic o in a typical com	of a				
wireless eq lical equipr nity to mea ystems' bas fully Autom unity test 1 60601-1-2 ance betwee	uipments ha nent and/or dical equipr ic safety ar hatic Digital evel in the :2014. The en RF wirele	ve being use systems are nent and/or nd essential Blood Press below table customer an	d in various sused. Whe systems, th performand sure Monito and meet to d/or user sl cations equi	healthcare n they are ne medical ce may be r has been the related hould help	communication: The device is int radiated therefor device can help minimum distan equipment (trans the maximum ou	s equipment and ended for use ir re disturbances prevent electror ce between port smitters) and th ttput power of th	I the device an electromag are controlled. nagnetic interf able and mobil e device as reco ne communicat	netic environment in which The customer or the user of the erence by maintaining a te RF communications ommended below, according to ions equipment.
Service	Modulation	Maximum power (W)	Distance (m)	Immunity test level (V/m)	Rated maximum output power of transmitter	80 MHz t	o 800 MHz	g to frequency of transmitter m 800 MHz to 2.7 GHz
	0 modulation 18Hz	1.8	0.3	27	W 0.01	- <u>1</u> 0		0.23
LTE Band	Pulse modulation	0.2	0.3	9	0.1	0.3	38	0.73
GSM 800/900 TETRA 800, iDEN 820, CDMA 850.	, Pulse modulation	2	0.3	28	10 100	12		7.3 23
GSM 1800; CDMA 1900;	Pulse	2	0.3	28	recommended se equation applica	paration distan ble to the frequ	ce d in metres (ency of the trar	(m) can be estimated using the nsmitter, where P is the maximum
		m 2	0.3	28			z, the separatio	on distance for the higher
WLAN 802.11 a/n	Pulse modulation 217Hz	0.2	0.3	9	NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.			
es		1	1	51	Additional N	otes		(
result in imp should be ob LE RF comm nal antennas Automatic Di R. Otherwis refer to the s susure accur press and ho. CD screen is splays the dc capacity, cali unanual pres then the diff ressure gauguracy. or infants or re monitor ar nent systems	rroper operati served to veri unications eq) should be us igital Blood P e, degradation software eval acy: Id the * STAR full, release puble zero, th ibrated standa momanomete ssure can be a ference betwe ge can be comp individuals w e reusable for designed to n sand adults in	on. If such us ify that they ar uipment (inclusion) ed no closer ti- ressure Monin on of the perfor- uation report, uation report, the "START/ e bloodpressur through the e pplied to the ce- en the reading- ne and ressure gas ho cannot exp e clinical and h neasure the sy dividual by u	e is necessary re operating r uding periph han 30 cm (1 tor, including mance of thill , and the file of ton, and then file of ton, and then file of ton, and then file of ton, and then STOP" button re meter is in auge and mar sleeve interfa ffective disp go fite sphysy ode can be use press their int nome use and sistoic and dir sing a non-in	r, this equipment iormally. erals such as 2 inches) to any g cables specified sequipment code is install h. static state. Jual pressure cc of the lay range of the gmomanometer ed to verify entions. are non-invasive astolic blood vasive technique	 9. ESSENTIÁL PERI Pressure calibration of the method described accuracy. If the accuracy deviai 10. Mechanical streng during the EXPECTE 11. Do not place the b child swallows or twi 12. The cuff and the c and do not contain all use. 13. Warning: Non-professionals dd measurement is not a 14. Warning: Do not expose the equ equipment. 15. Warning: This device is not use 16. Clean: The equipment can be instructions 17. Warning: Do not use a damageci 	[©] ORMANCE Mi will be carried on tin the section "' tion is large, ple- gth and resistanc D SERVICE LII lood pressure m ne around his ne asse of the blood dergenic or harm o not modify the ccurate. aipment for a lor d for children ar e cleaned by lay	aintenance advi at when this pro Verify Manomet ase contact the r e to heatThe res FE of the ME EC onitor and cuff a vek. pressure monit(ful materials.Pl equipment, other ag time, otherwi ad pets operator accord	ce: duct leaves the factory. Patients c ter Pressure Accuracy" to verify tl manufacturer to recalibration. sistance to heat will be retained by QUIPMENT. at will. It will cause asphyxiation or have been tested for biocompat lease stop using it if allergy occurs erwise it will make the equipment ise it will reduce the performance ling to the cleaning procedures in
	aration of n wireless eq ical equipy nity to me- stems' bas service TETRA 40 or service TETRA 40 or service Se	aration of manufacturer wireless equipments ha ical equipment and/or rity to medical equips stems' basics safety an ully Automatic Digital nity test level in the 60601-1-2:2014. The nece between RF wireled nad/or systems as rec Service Modulation TETRA 400 Puise TETRA 400 Puise and control and the rest-off and	aration of manufacturer-electromag wireless equipments have being use ical equipment and/or systems are risty to medical equipment and/or stems' basic safety and essential ully Automatic Digital Blood Press. unity test level in the below table 60601-1-2:2014. The customer an ince between RF wireless communit and/or systems as recommended 1 Service Modulation Maximum power (W) FETRA 400 modulation 1812 1.8 000 1.12:2014. The customer an ince between RF wireless communit and/or systems as recommended 1 Commender the systems as recommended 1 I TETRA 400 modulation 13.1 10 0.2 0.2 0.2 0.2 0.2 0.2 0.2 0.2 0.2 0.	aration of manufacturer-electromagnetic immur wireless equipments have being used in various ical equipment and/or systems are used. When hity to medical equipment and/or systems, th stems' basic safety and essential performance inty test level in the below table and meet to 60601-1-2:2014. The customer and/or user since between RF wireless communications equipation and/or systems as recommended below. Service Modulation Patie 0.3 Patie 0.3 IFETRA 400 Patie modulation 0.2 0.3 ISE 0.3 0.3 CSM 800/900, Patie 0.2 0.3 CSM 800/900, Patie 0.2 0.3 CSM 800/900, Patie 0.3 0.3 CSM 800/900, Patie 0.2 0.3 CSM 80	Service Modulation Maximum (W) Distance (m) Immunity test level (V/m) TETRA 400 Pulse (W) 1.8 0.3 27 Descent (V/m) Pulse (W) 0.3 27 Descent (V/m) 0.3 28 LTE (M) Pulse (M) 0.2 0.3 28 LTE (M) Pulse (M) 0.2 0.3 28 CSM 800%00 (COM 800% (COM 800%00) (COM 800%	aration of manufacturer-electromagnetic immunity wireless equipments have being used in various heather at the argent in the original Bood Pressure Monitor has been there and/or systems and/or systems, the machina distance of the below table and meet the related 60601-1.2:2014. The customer and/or argent should hedpend pressure Monitor has been and/or system as a recommended below.	The formulation of the prevent electron distance equipment and/or systems are used. When they are high the base, at Dey and cose Prese previous heathers the method equipment and/or systems are used. When they are high the base, at Dey and cose Prese previous the method electron the the related of the they are high to base, at Dey and cose Prese previous the method electron the they are high they have a they and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and cose Prese previous the method electron they are house at Dey and the Development (and they are house at Dey and the Development (and they are down and they are down at Development (and they are down at Development	Table 4 Table 4 Table 4 Table 4 Table 4 The device is intended separation distances between personal distances between

Additional Notes

20.Warning: This equipment is used outside the specified environment, may damage the equipment, and may be inaccurate measurement.

21.ME equipment not intended for use in conjunction with flammable agents "ME equipment not intended for use in oxygen rich environment'

X

Correct Disposal of This Product (Waste Electrical & Electronic Equipment)

This marking shown on the product indicates that it should not be disposed with other household waste at the end of its life. To prevent potential harm to the environment or to human health, please separate this product from other types of wastes and recycle it responsibly. When disposing this type of product, contact the retailer where product was purchased or contact your local government office for details regarding how this item can be disposed in an environmentally safe recycling center. Business users should contact their supplier and check the terms and conditions of the purchasing agreement. This product is hould not be mixed with other commercial wastes for disposal. This product is free of hazardous materials.

Safety Notice(Additional)

Federla Commulcation Commission (FCC) Interference Statement

 This device complies with part 15 of the FCC Rules. Operation is subject to the condition that this device does not cause harmful interference.
 This device is verified to comply with part 15 of the FCC Rules for use with cable television service. service. 3. This device complies with part 15 of the FCC Rules. Operation is subject to the following two

3. This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions: (1) This device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation. Please note that changes or modifications not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment. 4. This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to part 15 of the FCC Rules. These limits are designed to provide treasonable protection against harmful interference in a residential installation. This equipment generates, uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarante th interference will not occur in a particular installation. If this equipment does cause harmful interference to radio reception, which can be determined by turning the equipment off and on, the user is accouraged to try to correct the interference by one or more of the following measures: —Recorient or relocate the receiving antenna.

-Reorient or relocate the receiving antenna.

Recordent or recordent the receiving antennas.
 Increase the separation between the equipment and receiver.
 Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

is connected. —Consult the dealer or an experienced radio/TV technician for help. 5. This equipment complies with radio frequency exposure limits set forth by the FCC for an uncontrolled environment. 6. This device must not be co-located or operating in conjunction with any other antenna or transmitter.

7.Essential performance:

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Electrosurgery interference recovery	Refer 202.6.2.101	IEC 80601-2-30
Limits of the error of the manometer	Refer 202.12.1.102	IEC 80601-2-30
Reproducibility of the BLOOD PRESSURE DETERMINATION	Refer 201.12.1.107	IEC 80601-2-30