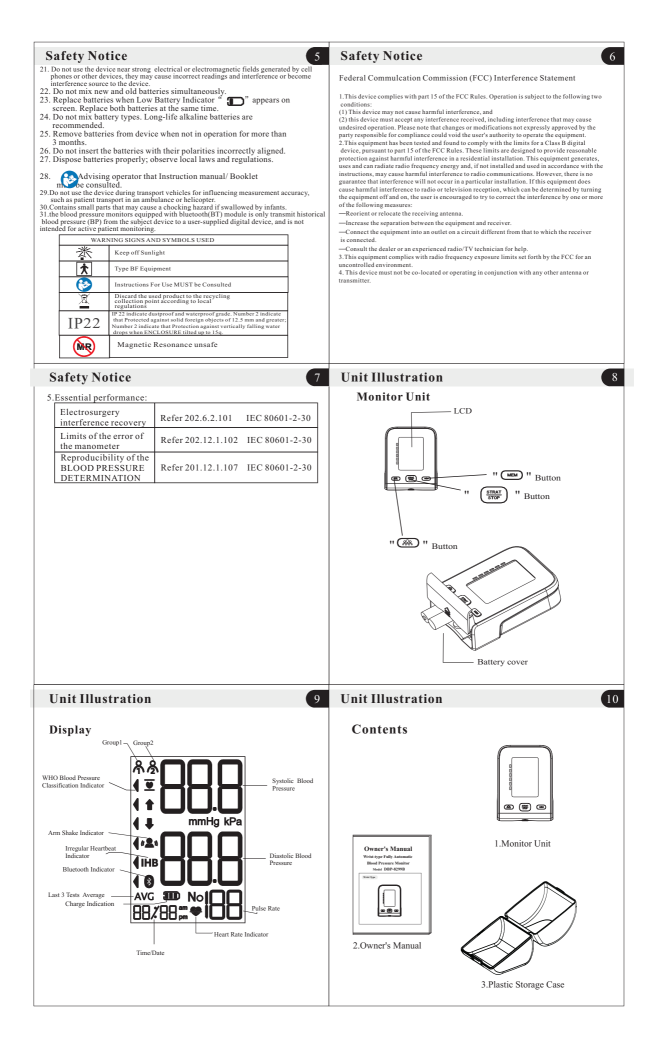
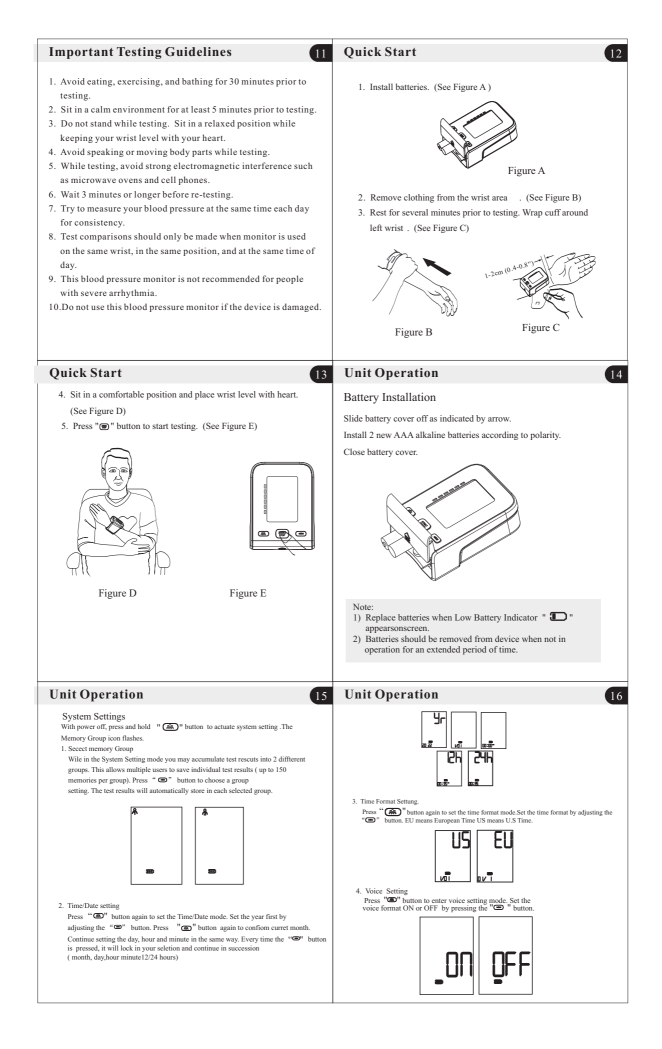
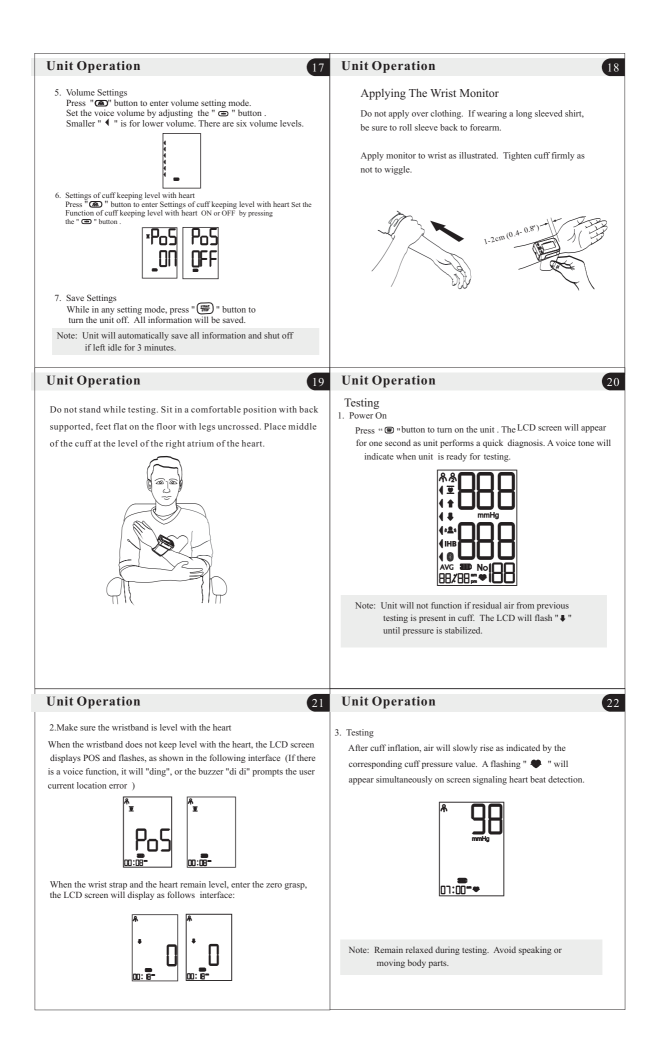
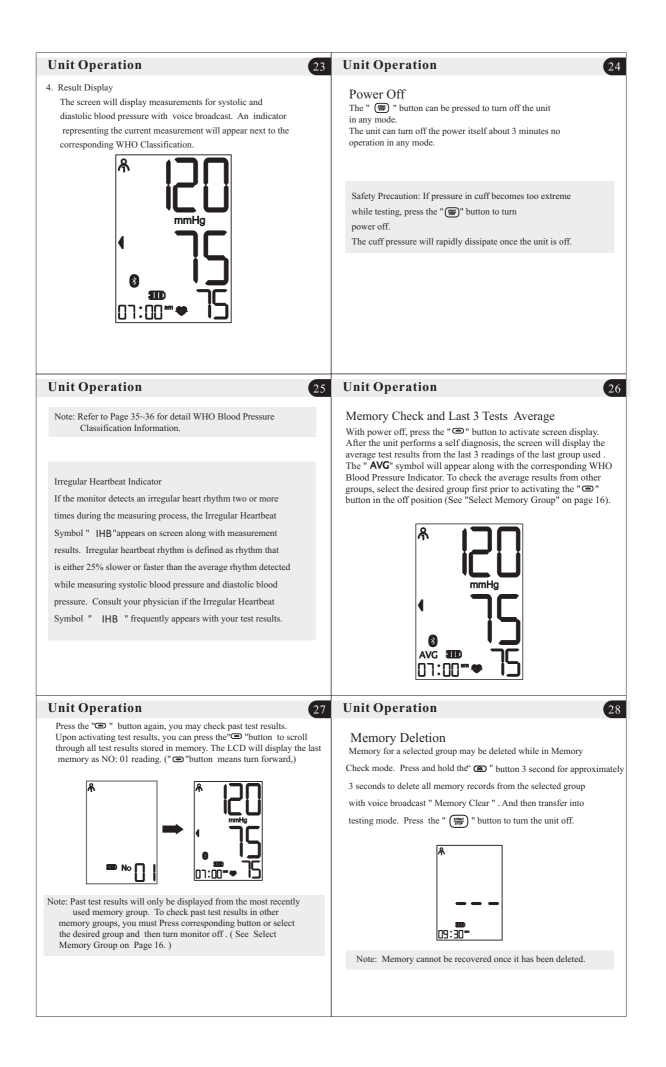


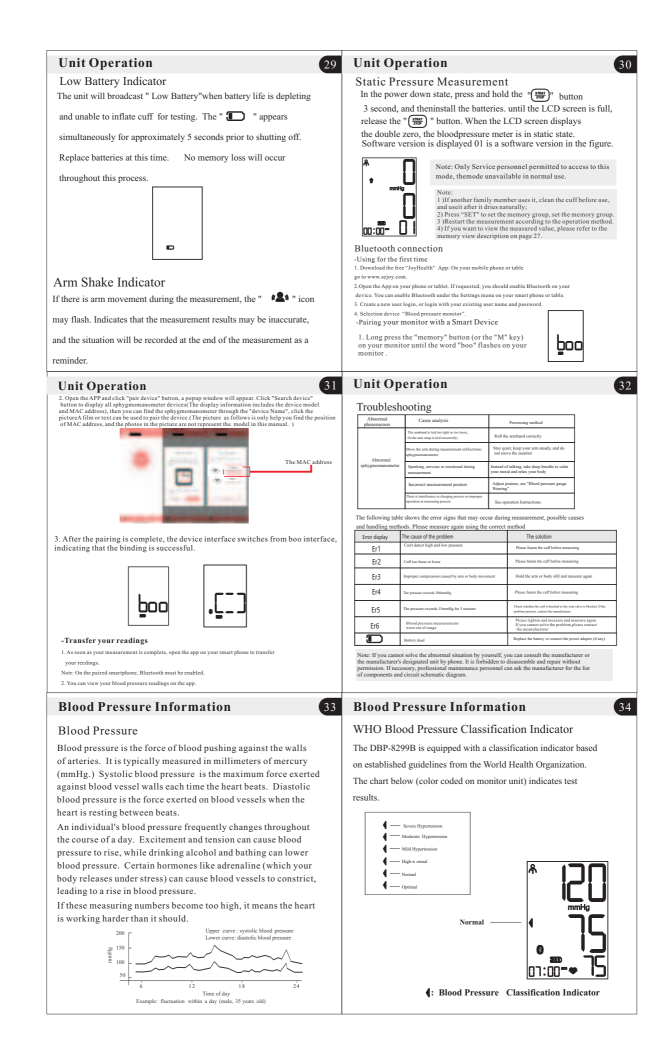
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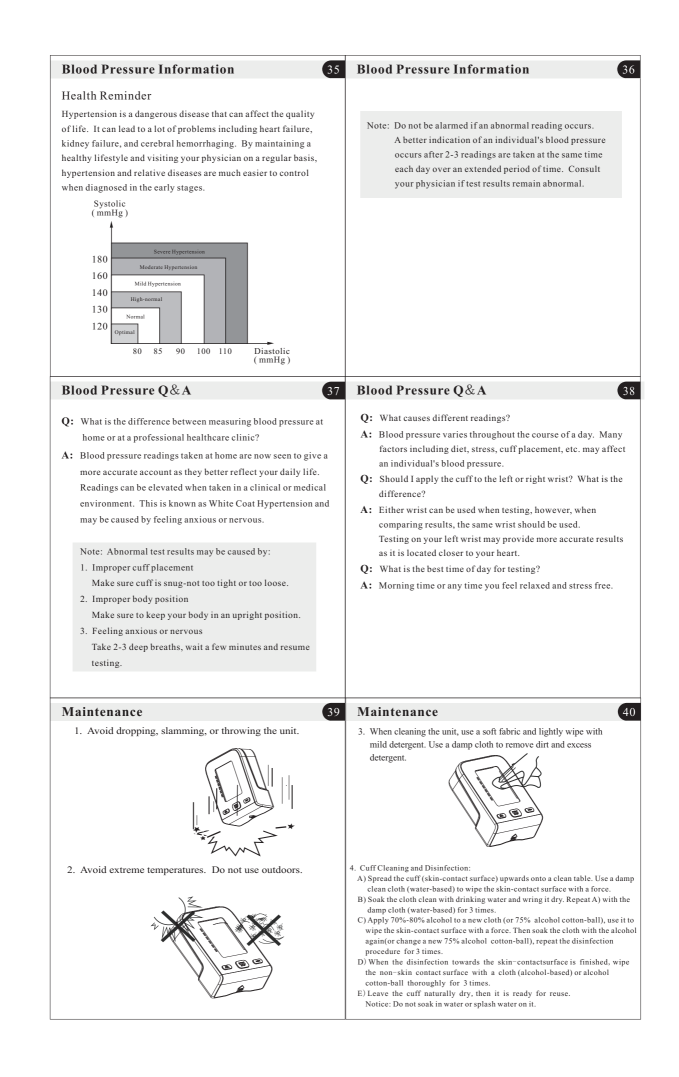












Vrist-type Fully Aut lood Pressure Mon PBP-8299B	tion for an extended tion for an extended omatic itor Size:45mm x 30.5mm(1.77" x 1.20") od 60mmHg~260mmHg 40mmHg~200mmHg ± 3mmHg 30~180 Beats/Minute	<ul> <li>2 years.</li> <li>9. Expected service liper day.</li> <li>10.No service and ma only be performed maintenance requibe provided.</li> </ul>	the performation of the performance with the performance with the performance with the performance with the performance of	ance should be checked even nately three years at 10 terminately three years at 10 terminately three years at 10 terminately three and mainteners onnel. Service and dir, technical support will there is Size AAA y 2 months at 3 tests per day (3.21 oz) (Excluding Battery)			
lood Pressure Mon DBP-8299B CD Digital Display Socillometric Metho vstolic Pressure astolic Pressure essure essure ulse	tomatic itor Size:45mm x 30.5mm(1.77" x 1.20") ad 60mmHg~260mmHg 40mmHg~200mmHg 0mmHg~299mmHg ±3mmHg 30~180 Beats/Minute	only be performed maintenance requibe provided.         3       Specifications         Function       Power Source         Battery Life       Unit Weight	by service per re parts, repa Voice Backlight Bluetooth 2 Alkaline Bat Approximatel Approx 91g Approx. 82.31	ersonnel. Service and ir, technical support will tteries Size AAA y 2 months at 3 tests per day			
lood Pressure Mon DBP-8299B CD Digital Display Socillometric Metho vstolic Pressure astolic Pressure essure essure ulse	tomatic itor Size:45mm x 30.5mm(1.77" x 1.20") ad 60mmHg~260mmHg 40mmHg~200mmHg 0mmHg~299mmHg ±3mmHg 30~180 Beats/Minute	Function Power Source Battery Life Unit Weight	Backlight Bluetooth 2 Alkaline Bat Approximatel Approx 91g Approx. 82.30	y 2 months at 3 tests per day			
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essure ulse	± 3mmHg 30 ~ 180 Beats/Minute	Unit Dimensions					
ulse	30 ~ 180 Beats/Minute		1 VI. 44 X 4.0	mm×63.8mm×27.5mm(L x W x "x 1.08")			
1150		Cuff Circumference	Fits wrist circ	umference 13.5-21.5 cm(5.3"-8			
tomatic Pressurizat	±5%		Temperature	$10^{\circ}C \sim 40^{\circ}C (50^{\circ}F \sim 104^{\circ}F)$			
	Fow Groups with Date and Time	Operating Environment	Humidity	15%~93%RH			
	*		Pressure	800hPa~1060hPa			
-		Storege Environment	Temperature	-25°C~55°C (-13°F~131°F)			
		Storage Environment	Humidity	≪93% RH			
st 3 Results Averag	je	Transport Environment	Temperature	-25°C~55°C (-13°F~131°F)			
w Battery Detectio	n	Transport Environment	Humidity	≪93% RH			
itomatic Power-Off	Ĩ						
	4	5 Specifications					
		Spacifications are subj	aat to ahange	without notice			
on Type GFSK		Safety Standard (included but not limited) :					
		1.IEC 80601-2-30, m	edical electri	cal equipment - part 2-3			
	(2400 <sup>-2</sup> 483. 5MHz)	performance of automa	performance of automated noninvasive sphygmomanometers.				
ndwidth 2.0 MHz		2.ISO 81060-2, non-in	2.ISO 81060-2, non-invasive sphygmomanometers - part 2:				
		3.AAMI / ANSI ES 60 C1:2009/(R)2012 and	0601-1:2005 a2:2010/0	(R) 2012 and (r) 2012 (consolidated tex			
nal Powered Equipr	ment Type BF	4.AAMI/ANSI/IEC 60	ial performan 601-1-2, M	nce edical Electrical Equipme			
H s v it	IO Classification 1 t 3 Results Average v Battery Detectio omatic Power-Off 5.0.1 BT frequency 2.4GHz in 0 dBi 2.0 MF	A Type GFSK 5.0.1 BT Signal mode frequency 2. 4GHz (2400 <sup>°</sup> 2483. 5MHz) in 0 dBi 2.0 MHz	IO Classification Indicator       Storage Environment         t 3 Results Average       Transport Environment         v Battery Detection       Transport Environment         omatic Power-Off       45         s 50.1 BT Signal mode       Specifications are subj         in 0 dBi       2.0 MHz         al Powered Equipment Type BF       Image: Specification and Cardiovascular)         al Powered Equipment Type BF       Image: Specification and Cardiovascular)	gular Heartbeat Detection       Temperature         IO Classification Indicator       Storage Environment       Temperature         Humidity       Temperature       Humidity         Transport Environment       Temperature         Mumidity       Temperature         Mumidity       Temperature         Transport Environment       Humidity         Transport Environment       Humidity         Mumidity       Specifications are subject to change         Sol BT Signal mode       Specifications are subject to change         Safety Standard (included but not li       I.EC 80601–2–30, medical electri         particular requirements for the basis       Cardiovascular)         2.1SO 81060–2, non-invasive sphy       Clinical validation of automated me         3AAMI / ANSI / ES 60601–1: 2005       Cl: 2009/ (R) 2012 and, a2: 2010/         medical electrical equipment Type BF       Fart 1-2: General Requirements for basic safet         Sesential Performance - Collateral Disturbances Collateral Disturbances Collateral And SJ.EC 60601–1-11, medical electrical el			

Warra	nty					4	7 E	lectromag	gnetic C	ompati	bility I	nformation
late of pu properly o vill repai	rchase. I due to de r or repla	f the Bloo fective co ace it free	od Pressu omponent ly. The w	nteed for 2 re Monitor s or poor v varranty do itor due to	does not vorkmans ves not co	function hip, we ver	6 tr p i: t t t s	0601-1-2. The re able below. The d recautionary mea nstructions for us the device. Use of he device negativ	equirements a levice is an e asures with r se. Portable a f the unit in c vely and alte	re satisfied lectrical me egard to EM nd mobile H onjunction	under the c dical produ C which mu IF commun with non-ap magnetic co	rnational standard IEC onditions described in the ct and is subject to special st be published in the ications equipment can aff proved accessories can aff ompatibility. The device er electrical equipment.
damages to your Blood Pressue Monitor due to improper handling. Please contact local retailer for details.								ended for use in	the electrom	agnetic envi	tic emissions ronment specified below. is used in such an environmen	
								Emissions test	c	ompliance	Electroma -guidance	gnetic environment
								Radiated emission		Group 1, ClassB	The device its internal emissions a likely to ca	uses RF energy only for function. Therefore, its re very low and are not use any interference in tronic equipment.
								Conducted emissi	on CISPR 11 P	i/A		
						Harmonic emissi IEC 61000-3-2	ons	J/A				
						Voltage fluctuati flicker emission	ons/					
								IEC 61000-3-3	~ I	I/A		
Electro	static correct(ESD) +	st level le 8 kV ± ntact cr 2 kV ± 4 kV ±	s 8 kV ontact 2 kV.±4 kV.	Electromagnetic -guidance Floors should be or ceramic tile. If covered with syn	wood, concrete floors are			IMMUNITY test	IEC 60601 test level	Compliar level	-guid Portabl equipm part of	e and mobile RF communications ent should be used no closer to an the device, including cables, than
Electro	static oge (ESD) 000-4-2	st level     le       8 kV     ±       ntact     cr       2 kV,±4 kV, ±     ±       8 kV, ±     ±	vel = 8 kV ontact = 2 kV,±4 kV,	-guidance Floors should be	wood, concrete floors are thetic material.						Portabl equipm part of the reco	e and mobile RF communications ent should be used no closer to an the device, including cables, than mmended separation distance ted from the equation applicable t uency of the transmitter.
Electro transie IEC 61	ostatic nt/burst 000-4-4 lin ± inp lin	2 kV for wer supply es 1 kV for out/output es 1 kV	N/A					Radiated RF EM fields IEC 61000-4-3	10 V/m80MH 2.7Ghz 80%A at 1kHz		%AM to 800 ! the may transmi transmi	mended seperation distance 80 M MHz 800 MHz to 2.7 Ghz where P imum output power rating of the tter in watts (W) according to the tter manufacturer and d is the tended separation distance in (m). Field strengths from fixed RI tters, as determined by an
Surge IEC 61	000-4-5 dif ± con	ferential	V/A								electro less tha frequen the vici followi	magnetic site survey, a should be n the compliance level in each icy range. Interference may occur nity of equipment marked with th ng symbol:
ons and variatio ower su put line	e dips, terrupti- upply in- 2000-4-11 cy/ cy/ cy/ cy/ cy/ cy/ cy/ cy/ cy/ cy/	95% dip in ') for 0.5 cle % UT 9% dip in ') for 5	ν/A					Conducted disturbances Induced by RF fields IEC 61000-4-6	3 V in 0.15 MHz- 80 MH 6 V in ISM and/or amateu radio bands between 0.15 MHz and 80 MHz 80 % AM at 1kHz	6 V in ISM	equipm part of recomm 5 MHz 4 10 800 1 15 15 15 15 15 15 15 15 15 15 15 15 15	e and mobile RF communications or thould be used no clear to an end thould be used no clear to an ended separation distance the deform the equation applicable usency of the transmitter. Will z800 MHz to 2.7 G che where x80 M MHz 800 MHz to 2.7 G che where immun output power rating of the inter manufacturer and d is the mended separation distance in (m). Field strengths from fixed 8 magnetic atte survey, a should be compliance level in each freque interference may occur in the interference may occur in the g symbol: $\frac{12}{10000000000000000000000000000000000$
(50/60) magnet IEC 610	Prequency Hz) ic field 30 000-4-8	60Hz o	0 A/m; 50Hz r 60Hz	Power frequency should be at leve typical location i rcial or hospital	ls charactertic n a typical com environment.	of a me-						
able 3	0			r-electromag			Т	lectromag able 4 Recommended so	-	_		Information
cations w ed in clo juipment fected. A sted with quiremen eep a mini	where medi ose proxim and/or sy rm-type Fu the immu- ts of IEC mum dista	ical equipm nity to med stems' basi ally Autom nity test le 60601-1-2: nce betwee	ent and/or ical equips c safety as atic Digital evel in the 2014. The n RF wirele	ve being use systems are nent and/or nd essential Blood Press below table customer an ess communi commended l	used. Who systems, th performan sure Monito and meet d/or user s cations equ	en they are ne medical ce may be or has been the related hould help		communications The device is inte radiated therefore device can help pr minimum distanc	equipment a ended for use e disturbance revent electr e between po mitters) and	nd the devic in an electr s are contro omagnetic i ortable and r the device a	e omagnetic o illed. The cu nterference nobile RF c s recommen	environment in which astomer or the user of the by maintaining a ommunications ided below, according to
Test quency MHz)	Band (MHz)	Service	Modulation	Maximum	Distance (m)	Immunity test level (V/m)	01	atput power of ansmitter	1.5			800 MHz to 2.7 GHz
385 450	380-390 430-470	TETRA 400 GMRS 460 FPS 460	modulation 18Hz ± 5 kHz deviation	1.8	0.3	27		W 0.01		0.12		0.23
710 745	704-787	LTE	Pulse	0.2	0.3	9		0.1	(	0.38		0.73
780	/04-787	Band 13, 17	modulation 217Hz	n 0.2	0.0	, ,		1		1.2 3.8		2.3 7.3
810 870	800-960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse modulation 18Hz	n 2	0.3	28		100		12		23
930 1720 1845	1700-1990	LTE Band 5 GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse modulatior 217Hz	1 <u>2</u>	0.3	28		recommended sep equation applicat	paration dist ble to the free	ance d in me juency of th	tres (m) can e transmitte	t listed above, the a be estimated using the er, where P is the maximum coording to the transmitter

0.3

0.3

2

0.2

28

9

2450

5240

5500

5785

2400-2570

5100-5800

actooth,WLAN .11 b/g/n,RFID 50,LTE Band 7

WLAN 802.11 a/n Pulse modulatio 217Hz

Pulse nodulation 217Hz NOTE1 At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

## Additional Notes

## Important Instructions Before Use

WARNING: Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally. 2.WARNING: PORTABLE RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of Wrist-type Fully Automatic Digital Blood Pressure Monitor, including cables specified by the MANUFACTURER. Otherwise, degradation of the performance of this equipment could result.

3. The software identifier refer to the software verification and validation report, and the file code is JYRJ200930001.

sphygmomanometer, and manual pressure can be applied to the effective display range of the sphygmomanometer, and then the difference between the reading of the sphygmomanometer and that of the standard pressure gauge can be compared. This mode can be used to verify manometer pressure accuracy.

5.Contraindications:

Product is not intended for infants or individuals who cannot express their intentions 6.Intended Use

The digital blood pressure monitor are reusable for clinical and home use and are non-invasiv blood pressure measurement systems designed to measure the systolic and diastolic blood pressure and pulse rate of adolescents and adults individual by using a non-invasive techniqu , which is a well-known technique in the market called the "oscillometric method". it can measure the systolic blood pressure, diastolic blood pressure and pulse rated on

up-arm, and the device is reusable for clinical or home use. 7.The patient is the operator: the PATIENT is an intended OPERATOR. the PATIENT Do not carry out other maintenance operations except to replace the battery

**Additional Notes** 

# 55

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



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

This marking shown on the product indicates that it should not be disposed with other 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 should not be mixed with other commercial wastes for disposal. This product is free of hazardous materials.

### **Additional Notes**

#### 8 WARNING

53

Do not modify this equipment without authorization of the manufacturer 9. ESSENTIAL PERFORMANCE Maintenance advice:

Pressure calibration will be carried out when this product leaves the factory. Patients can use the method described in the section "Verify Manometer Pressure Accuracy" to verify the

accuracy. If the accuracy deviation is large, please contact the manufacturer to recalibration

In the actual y deviation is happened to hear the resistance to hear will be retained by device during the EXPECTED SERVICE LIFE of the ME EQUIPMENT. 11.Do not place the blood pressure monitor and cuff at will. It will cause asphysiation if the child swallows or twine around his neck.

2. The cuff and the case of the blood pressure monitor have been tested for biocompatibility and do not contain allergenic or harmful materials. Please stop using it if allergy occurs during use

13. Warning: Non-professionals do not modify the equipment, otherwise it will make the equipment measurement is not accurate.

14.Warning:

Do not expose the equipment for a long time, otherwise it will reduce the performance of the equipment. 15.Warning:

This device is not used for children and pets 16.Clean:

The equipment can be cleaned by lay operator according to the cleaning procedures in the instructions

17. Warning: Do not use a damaged cuff for blood pressure measurement. 18. Warning:

When measuring with the cuff, if the tester feels seriously uncomfortable, press the button of the blood pressure monitor to deflate the cuff, or remove the cuff directly from the arm 19.Warning:

If an unexpected reading occurs, the operator can take several more measurements and consult a doctor.

