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Product Description	Wrist-Type Fully Automatic			Function	Buzzer			
*	-	d Pressure Monitor		Power Server	Backlight	m. (2 m. 18 ar 1 b)		
fodel Display	DBP-8276H LCD Digital D	isplay Size:34mm×34.6mm (1.34"	x 1.36")	Power Source	lithium battery (3.7v 180mAh)			
feasurement Method	Oscillometric Method			Battery run time	500 measurements			
	Systolic Pres	sure 60mmHg~260mmHg		Charge time Unit Weight	Approx. 122g (4.300z.) (Excluding Batter			
	Diastolic Pres	sure 30mmHg~20ommHg		Unit Dimensions	Approx. 62mm×55.2mm×19mm(L x W x H) (2.44 " x 2.17"x 0.75")			
Measurement Range	Pressure	0mmHg~299mmHg		Cuff Circumference	Fits wrist circumference 13.5-21.5 cm(5.3			
measurement Kange	Pressure	±3mmHg			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%			Pressure	800hPa~1060hPa		
essurization	Automatic Pr	essurization			Temperature	-25℃~55℃ (-13°F~131°F)		
emory	2x60 Memories in Tow Groups with Date and Time			Storage Environment	Humidity	≪93% RH		
	Irregular Heartbeat Detection			Storage Environment	Temperature	-25℃~55℃ (-13℉~131℉)		
	WHO Classif	ication Indicator		Transport Environment	Humidity	≤93% RH		
unction	Last 3 Result	s Average						
	Low Battery	Detection						
	Automatic Po	wer-Off						
pecifications	\$		43 S	pecifications				
	Modulation Type	GFSK	Sı	pecifications are subj	ject to chans	ge without notice.		
	Version					-		
	Operation	5.0.1 BT Signal mode	ar	nd bears the CE mark'	"CE 0197".	plies with the European regula This blood pressure monitor a		
Bluetooth	frequency	2. 4GHz(2400 [~] 2483. 5MHz		omplies with mainly f ncluded but not limit		andards		
	Antenna gain	0.5 dBi	- Sa	afety standard:	í.	ipment part 1: General		
	Bandwidth	2.0 MHz	re	quirements for safety		apment part 1. Ocherai		
gress Protection Rating	IP 22		El			Equipment Part 1-2: Genera		
assification	Internal Powe	ered Equipment Type BF 🚺	Sa		erformance	Collateral Standard:		
attery Shelf life	60 months		R	Electromagnetic Disturbances – Requirements And Tests. Performance standards:				
emperature	1		re		asic safety a	l equipment – Part 2-30: Part: nd essential performance of manometers		
			re sy IS	N 1060-3 Non-invasi quirements forelectro ystems.	ve sphygmo omechanica sive sphygi	manometers - Supplementary 1 blood pressure measuring nomanometers - part 2: clinic		
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The device is in The customer or ent.	the user of the dev	he electromagneti vice should assure	that it is used ir	1 such an enviror	m-	ent. IMMUNITY test	IEC 60601	Compliance	Electromagnetic environmen
Electrostatic	± 8 kV	± 8 kV contact	-guidance Floors should be or ceramic tile.	e wood, concrete	_		test level	level	-guidance Portable and mobile RF communicati equipment should be used no closer to part of the device, including cables, the the recommended separation distance
discharge (ESE IEC 61000-4-2 Electrostatic	± 15 kV air ± 2 kV for power supply	±8 kV,	he relative hum t least 30 %.	nthetic material, idity should be	-	Radiated RF EM	80MHz-2.7	3V/m or 10 V/m 80MHz-2.7	the recommended separation distance calculated from the equation applicat the frequency of the transmitter. Recommended seperation distance 8(to 800 MHz 800 MHz to 2.7 Ghz whee the maximum output power rating of transmitter in watts (W) according to
transient/burst IEC 61000-4-4	lines ± 1 kV for input/output lines ± 1 kV differential	N/A			_	IEC 61000-4-3	Ghz 80%AM at 1kHz	Ghz 80%AM at 1kHz	transmitter manufacturer and d is the recommended separation distance in metres (m). Field strengths from fixed
Surge IEC 61000-4-5	mode ± 2 kV common mode < 5% UT	N/A			_				electromagnetic site survey, a should less than the compliance level in each frequency range. Interforence may oc the vicinity of equipment marked with following symbol: we Portable and mobile RF communicati
Voltage dips, short interrupti ons and voltage variations on p- ower supply in- put lines IEC 61000-4-1	cycle 70% UT (30% dip in UT) for 25	N/A				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	equipment should be used no closer t part of the device, including cables, tr ecommended separation distance the frequency of the transmitter. Recommended seperation distance 80 transmitter in watts (W) according to transmitter in according to the transmitter matters (m). Field strengths from five frame, Interference may occur in the vicinity of equipment marked with th following symbols: $\frac{1}{2}$
Power frequenc (50/60 Hz) magnetic field IEC 61000-4-8		30 A/m; 50Hz	should be at lev	y magnetic field els charactertic o in a typical com environment.	fa				
e 3 uidance and c adays, many tions where n in close pro- pment and/or ted. Arm-typ d with the in irements of l	RF wireless e nedical equiporimity to m systems' ba e Fully Auto nmunity test EC 60601-1-	manufacturer quipments ha ment and/or edical equip tsic safety an matic Digital level in the 2:2014. The	-electromag ve being use systems are nent and/or nd essential Blood Pres below table customer an	gnetic immur ed in various e used. Whe systems, th performanc sure Monito e and meet t nd/or user sl	healthcare n they are e medical e may be r has been he related nould help	Table 4 Recommendeds communications The device is int radiated therefor device can help j minimum distan	eparation dista s equipment and ended for use in re disturbances prevent electron ce between por	nces between p d the device n an electromag are controlled. magnetic interf table and mobi	ility Information ortable and mobile RF metic environmer in which The customer or the user of th erence by maintaining a le RF communications ommended below, according t
nedical equip	ment and/or	een RF wirele systems as rec	ommended Maximum	below.	Immunity	the maximum ou Rated maximum			ions equipment. g to frequency of transmitter
Ban (MH 85 380-3	z) Service	Pulse 00 modulation	power (W)	Distance (m) 0.3	test level (V/m) 27	output power of transmitter W	80 MHz $d = \left[\frac{3.5}{E_1}\right]$	to 800 MHz]√P	800 MHz to 2.7 GHz $d = \left[\frac{7}{E_1}\right]\sqrt{P}$
50 430-4	70 GMRS 460 FRS 460	18Hz FM ± 5 kHz deviation 1 kHz sine	2	0.3	28	0.01	0.		0.23
10 704-7	87 LTE Band 13, 17	Pulse modulation	0.2	0.3	9	0.1	0.:		0.73
80	GSM 800/9	21,112				10 100	3.		7.3
70 800-9 30 720 345 1700-1	GSM 1800 CDMA 859 LTE Band	; [;] [;] [;] [;] [;] [;] [;] [;] [;] [;]	2	0.3	28	For transmitters recommended se equation applica output power rat	rated at a maxim paration distant ble to the frequ	mum output po ice d in metres ency of the trai	wer not listed above, the (m) can be estimated using the ismitter, where P is the maxim (W) according to the transmit
970	LTE Band 1 4, 25; UM					NOTE1 At 80 M frequency range		z, the separatio	on distance for the higher
450 2400-2 240	570 Bluetooth,WL 802.11 b/g/n,RI 2450,LTE Band	AN, TD Pulse modulatio 17 217Hz	n 2	0.3	28	NOTE2 These gr	uidelines may n		situations. Electromagnetic
500 5100-5 785	800 WLAN 802.11 a/n	Pulse modulation 217Hz	0.2	0.3	9	propagation is a and people.	ffected by absor	rption and refle	ction from structures, objects
litional N	otes				51	Additional N	otes		
d because it cc events of the comparison of the events of the comparison of the comparison of the standa not intered the standa not intered the standa eter pressure is and intered raindications: t is not intend- ided Use gital blood pre- pressure measus can and pulse raison of the events of the standa standard the standa of the standa of the resource of the standa of the standa of the standa of the standard the standard standard the standard the standard the standard standard the standard	uld result in in ent should be or ABLE RF com ternal antenna URER. Otherw fier refer to the pressure accu te, press and h e LCD screen displays the tas capacity, cz ted to the sphy and manual pr and then the d of pressure gat we curacy. ed for infants o ssure monitor a rement system	bserved to veri munications eq s) should be us Digital Blood P ise, degradatio e software eval racy: old the "STAR is full, release double zero, th librated stand gmomanomete essure can be a r individuals w re reusable for is designed to r ts and adults in	on. If such us fy that they a uipment (inc. ed no closer i ressure Mon. of the perfo- uation report uation report T/STOP" but the "START; e bloodpress ir dpressure e through the pplied to the en the readin pared. This m ho cannot ex clinical and neasure the sp dividual by u	te is necessary re operating r luding periph than 30 cm (12 itor, including rmance of this , and the file of ton, and then STOP ⁺ buttor are meter is in auge and man sleeve interfa effective disp g of the sphys ode can be use	t, this equipment ormally. erals such as inches) to any c cables specific e equipment code is install t. static state. ual pressure cc of the lay range of the momanometer d to verify entions. are non-invasiv ustolic blood vasive technique	Pressure calibration 1 the method described accuracy. 1 If the accuracy devia 10. Mechanical streng during the EXPECTF 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 dc 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:	ORMANCE M will be carried o in the section " in the section " the section of the section" D SERVICE LI lood pressure m ne around his n ase of the blood dergenic or harm o not modify the ccurate. aipment for a lor d for children an e cleaned by lay	aintenance advi ut when this pro- Verify Manome ase contact the : ie to heatThe re: FE of the ME EF onitor and cuff: reck. pressure monite ful materials.Pl equipment, oth ng time, otherw nd pets operator accord	ce: duct leaves the factory. Patient ter Pressure Accuracy" to verify manufacturer to recalibration. istance to heat will be retained QUIPMENT. at will. It will cause asphyxiatio or have been tested for biocomp ease stop using it if allergy occ erwise it will make the equipme ise it will reduce the performan ling to the cleaning procedures

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'

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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 guarantee that 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 of fand on, the user is encouraged 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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7.Essential performance.		
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