

Arm Blood Pressure Monitor

Model:U807



Instruction Manual

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of wrong measurement -----

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▲ Your new digital blood pressure monitor uses the oscillometric

method of blood pressure measurement. This means the monitor

detects your blood's movement through your brachial artery and

converts the movements into a digital reading. An oscillometric

monitor does not need a stethoscope, so the monitor is simple to

▲ Intelligent inflation will reduce the uncomfortable feeling by

incorrect inflation, and shorten the measurement time, prolong

▲ 2x90 sets memory function each measurement result will be

displayed on the screen, and automatically stored. This unit has

index, could easy to check your classification index.

blood classification index, could easy to check your classification

▲ Please read the manual carefully before you use the unit, and

This product can't be used in patients who is with severe heart

insufficiency to avoid suffocation and death. This product is not

This automatic blood pressure monitor intends to measure the

upper arm. It's expected to be used at home or in the hospital,

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systolic pressure, diastolic pressure and pulse rate through

Introduction ----

Product Structure

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-- How to set -

Proper use of the unit

-- Common factors

Introduction

the cuff's usage lifetime.

CONTRAINDICATION

INTENDED USE

keep the manual well after using.

suitable for infants and children.

intended for people over 12 years old.

Safety Information

■ To assure the correct use of the product, basic safety measures should always be followed including the warning and the caution listed in the instruction manual:

Symbol descriptions

The following symbols may appear in this manual, on the label, on the device, or on its accessories. Some of the symbols represent standards and compliances associated with the

▲ WARNING: This alert identifies hazards that may cause serious personal injury or death.

minor personal injury, product damage or property damage.

★ Type BF applied part

Manufacturer

SN Specifies serial number

EC REP Authorized Representative in the European Community

DISPOSAL: Do not dispose of this product as unsorted municipal waste. Collection of such waste separately for special treatment is necessary.

Direct current

Follow instructions for use

Safety Information

⚠ Those who have arrhythmia, diabetes, blood circulation or apoplexy problem, please use under the physician's instruction.

⚠ Contact your physician for specific information about your blood pressure. Self diagnosis and treatment which use measured results may be dangerous. Follow the instructions of your physician or licensed healthcare

A Please place on a high place where children can't be

A No modification of this equipment is allowed.

testing must be conducted to ensure continued safe use of

The cuff hose around neck may cause the suffocation.

⚠ Please don't use a dilution agent, alcohol or petrol to clean the unit. Please don't hit heavily or fall down the product from a high place. Use the right cuff, otherwise it can not

case of device with rechargeable lithium battery).

⚠ Please avoid using in high radiant area in order to make your measuring data correctly.

⚠ Do not modify this equipment without authorization of the

⚠ If this equipment is modified, appropriate inspection and

⚠ The swallowing of small part like packaging bag, battery,battery cover and so on may cause the suffocation.

⚠ Do not replace or remove the battery from device (in the

 \triangle Do not use a cellular phone near the unit. It may result in

⚠ Do not use the equipment where flammable gas (such as anesthetic gas, oxygen or hydrogen) or flammable liquid (such as alcohol) are present.

M WARNING:

municipal waste. Use separate collection facilities. are disposed of in landfills or dumps, hazardous

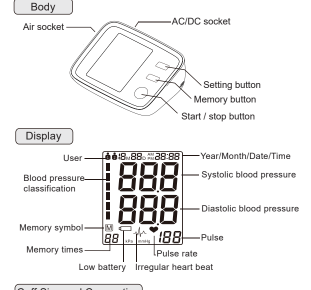
Classification

- 1. Internally powered equipment;
- 2. Type BF applied part;
- 3. Protection against ingress of water or particulate matter: IP21;
- 4. Not category AP / APG equipment;
- 5. Mode of operation: Intermittent operation.

⚠ The user must check that the equipment functions safely and see that it is in proper working condition before being used.

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Product Structure



Cuff Size and Connection

The accessories cuff is universal size for upper-arm circumference 22-42 cm use. The cuff is treated as the applied part. Insert the connector with cuff tube into the hole which is on the left side of the

device as pictured

change to any other branded cuff.)

Remove the battery cover from the battery compartment

the SIM card slot is on the right side (as picture shown). Insert SIM card into the slot in accordance with direction

O Do not replace or remove the battery from device.

When power on, the low battery symbol will be displayed once the

device starts. Plug in the device using the included USB cable to charge.

Take the USB cable and connect it to the charger (charge connector) or

to your USB port on your computer or power bank. Connect the end of

During charging when power off, the LCD will shows the blood

will be displayed once battery is full, as picture shown.

pressure bars on the left side from one bar to six bars in a continuous

8

The device is equipped with 1pc 3.7V 1000mAh

the round tip to the USB socket of device.

for about 70 times when charge

is full. Charge time is 4-6 hours.

Noted: the lithium battery could be used

Battery installation

SIM card installation

Battery installation

Noted:

rechargeable lithium battery.

LOW battery and charge

(Only provided cuff can be used, can not

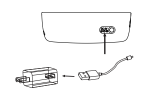
Setting mode

4. Insert the other side of the adapter into the outlet with 100-240V

5. To remove the AC adapter, disconnect the adapter plug from the outlet first and then disconnect the cord from the unit's socket.

Adapter technical features:

Output voltage: 5V±5% Max output current: At least 1A Output plug polarity: <+> inner External diameter: 5.5 mm 0.1mm Internal diameter: 2.1mm 0.1mm



Note:

- · When use AC adapter, the power of battery won't be consumed.
- · When suddenly stop during measurement (like the plug off from the outlet by carelessness), it must be reinserted the plug into the unit, and restart the measurement.

How to set

1. Start to set, Unit setting:

Setting mode

SPE OFF: close the speaking function. SPE ON: open the speaking function.

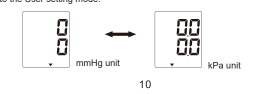
2. Speaking setting

3. User setting:

4. Year setting:

SIM card slot

Press the SET button when power off, 0 or 0.0 will be displayed, then the setting begin Continute to above step, the unit will be changed when press the MEM button each time. Press the SET button to confirm the unit, then it will enter into the User setting mode



Continute to above step, the screen will display SPE OFF or SPE ON, press the MEM button and it will change between SPE OFF and SPE ON, press the SET button to confirm the option. Following this, the device will enter

Continue to above step, the screen will display 🐧 or 🐧, press

button MEM ,it will be changed between 🛍 and 🛍 , press button SET

when you confirm the user, then it will enter into the year setting mode.

Continue to above step, the screen will display and flash 20XX, the last

digit of the year will increase 1 when press button MEM each time, you

could choose from 2020 to 2099. Press button SET when you confirm

the year, then it will enter into the month and date setting mode.

SPE

user 2

Proper Use of the Unit

Proper use of the unit

for 30 minutes before taking a measurement.

blood pressure changes even during the day.

Common factors of wrong measurement

• All efforts by the patient to support their arm can increase

• Make sure you are in a comfortable, relax position and do no

measurement. Use a cushion for support if necessary.

• If the arm artery lies lower or higher than the heart,a false

• A loose cuff or a exposed bladder causes false reading.

Consecutive blood pressure measurements should be

up in order to allow the accumulated blood to flow away.

With repeated measurements .blood accumulates in the arm

repeated after 1 minute pause or after the arm has been held

tactivate any of the muscles in the measurement arm during

Relax for about five to ten minutes prior to the measurement

All these factors will influence the measurement result.

• Remove any garment that fits closely to your upper arm. Always measure on the same arm(normally left)

Take measurement regularly at the same time of every day,as

Avoid eating, drinking alcohol, smoking, exercising and bathing

Fitting the Cuff

blood pressure

Note:

reading will be obtained.

• Only use clinically approved cuffs!

which can lead to false reading.

1.) Place the cuff flat on the table with the velcro side down. Pass the end of the cuff through the metal loop so that a circle is formed. The velcro closer will now be facing outwards (ignore this step if the cuff has already been prepared).

2.) Push the cuff over the left upper arm so that the tube points in the direction of the lower arm.

3.) Wrap the cuff on the arm as illustrated. Make certain

that the lower edge of the cuff lies approximately 2 to 3 cm above the elbow and the rubber tube leaves the cuff on the inner side of the arm.

4.) Tighten the free end of the cuff and close the cuff by affixing the velcro.

5.) The cuff should be snug on your upper arm so

That you can fit 2 fingers between the cuff and your upper arm. Any piece of clothing restricts the arm and should be removed.

6.) Secure the cuff with the velcro closer in such a way that it lies comfortably and not too tight. Lay your arm on a table (palm upwards) so that the cuff is at the same height as the heart. Do not bend the

If it is not possible to fit the cuff to your left arm, it can also be placed on the right. However, all measurements should be made using the same arm

150

6 10×20; 8:00

145

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End

Proper use of the unit

Measuring Procedure:

After the cuff has been appropriately positioned, the measurement can begin as follows: 1. Press the START/STOP button, all symbols appear on the display, then the pump begins to inflate the cuff, the rising pressure in the cuff is shown on the display.

2. After the suitable pressure has been reached, the pump stops and the pressure gradually falls. The cuff pressure is displayed. In case that the inflation is not sufficient, the device automatically re-inflates to a higher pressure.

. When the device detects the signal, the heart symbol • on the display starts to flash. When the measurement has been completed. the systolic, diastolic and pulse rate will

appear on the display. Remark: Once finish the meas device it will upload the dada to the background server automatically with showing two rotating signal bar in the top right corner of LCD (as Pic 1). After uploading the data successfully, the LCD will show End (as Pic 2) and then the device

switches off immediately. If fail to upload the data Note: The symbol $\sqrt{}$ will be displayed along with the reading if the irregular

heartbeat is detected during the measurement Discontinuing a measurement If it is necessary to interrupt a blood pressure measurement for any reason

(eg. the patient feels unwell) the START/STOP button can be pressed at any time. The device immediately decrease the cuff pressure automatically. Memory-recall of measurements

This blood pressure monitor automatically stores 2x90 sets measurements value, the oldest record will be replaced by the latest measurement value

be shown, press the button MEM again, the last measurement value will be shown, as well as subsequent measurements can be display one after the other by pressing the button MEM each time.

Safety Information

Do not dispose of electrical appliances as unsorted Contact you local government for information regarding the collection systems available. If electrical appliances substances can leak into the groundwater and get into the food chain, damaging your health and well-being.

Battery installation

If the device under setting mode and battery charging in the same time, LCD can show the same as above, e.g. user setting



▲ WARNING:

Dispose of the battery in accordance with all federal, state and local laws. To avoid fire and explosion hazard, do not burn or incinerate the battery

1. When optional AC adapter is in use, it should comply with the

Adapter Usage

requirement of IEC 60601-1:2005. Furthermore, all configurations shall comply with the requirements for medical electrical systems (see IEC 60601-1-1 or clause 16 of the 3Ed. of IEC 60601-1. respectively). Anybody connecting additional equipment to medical electrical equipment configures a medical system and is therefore responsible that the system complies with the requirements for medical electrical systems. Attention is drawn to the fact that local laws take priority over the above mentioned requirements. If in doubt, consult your local representative or the technical service department. 2. When using AC power, to avoid possible damage to the monitor, use only the exclusive AC adapter that can be purchased from authorized dealers. Other adapters may vary in output voltage and polarities. 3. Insert the adapter plug into the hole on the backside of the unit

as pictured 9

Setting mode

5. Month and date setting

Continue to above step, the screen will display xxMxxD and xxxx, and keep flashing on month, the digit will increase 1 when press button MEM each time, you could choose from 1 to 12. Press button SET when you confirm the month, then it will set the date. Same as the month setting, each time you press button MEM, the digit will keep changing from 01 to 31. Press button SET when you confirm the date, then it will enter into the time setting mode.



keep flashing on the digits of hour, the digit will increase 1 when press button MEM each time, you could choose from 0 to 23. Press button SET when you confirm the hour, then the digits of minute start to flash, same as the hour setting, each time you press button MEM the digits will keep changing from 00 to 59. Press button SET when you confirm the minute, then the total setting mode is completed.



Pre-measurement

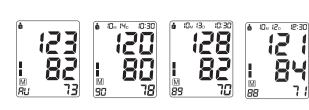
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Continue to above step, the screen will display xxMxxD and xx:xx, and

LCD will display Err.

when more than 90 sets each user. Read memory record Press the button MEM when power off, the latest 3 times average value will

About blood pressure



Memory -clear of measurements

If you are sure that you want to permanently remove all stored memories. Press the button SET for 9 times until CL appears when power off, press the START/STOP button ,CL will flash for 3 times to clear all the memories. After this press button MEM, M and "no" will be shown on the display which mean that no memory in store.

Check IMEI details:

Hold and press the MEM button for 3 seconds when power off, released the button and wait for 5 seconds, then the LCD will display the device IMEI number. And press the START/STOP button to exit.



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Exceptional Situation

Error Indicators

THE IONO	wing symbol will app	
Symbol	Cause	Correction
E- 1	Weak signal or pressure change	Wrap the cuff properly.
suddenly		Remeasure with correct way.
E-2	External strong	When near cell phone or other high radiant device, the measurement will be failed.
	disturbance	Keep quite and no chatting when measure.
	It appears arror	Wrap the cuff properly.
E-3	process of	Make sure the air plug is properly inserted into the unit.
	inflating	Remeasure.
٤-5	Abnormal blood pressure Repeat the measurement after relax for 30 mins, if unusual readings for 3 times, please contact your do	
	Low battery	Connect the USB cable and charger to charge

■ The following symbol will appear on the display when measuring abnormal

Trouble removal

y power Charge the battery insert Insert into the air socket tightly throken or leak Change a new cuff
broken er leek Change a new cuff
broken or leak Change a new cuff
e arm when inflate Keep the body peaceful
when measured Keep quite when measure
wrap too loose Wrap the cuff tightly

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Warranty Information

- The intended use: the unit is intended to be used by adults at home or medical center to measure blood pressure and pulse rate from the
- The device meets the requirements of IEC 80601-2-30 Part 2 for non-invasive blood pressure monitors.
- Blood pressure measurements determined with this device are equivalent to those obtained by a trained observer using the cuff/stethoscope auscultatory method, within the limits prescribed by the American National Standard, manual, electronic, or automated sphygmomanometers
- The risk of patient and user can be lowered to acceptable level.

Warranty Information

- The unit is guaranteed to be free of defects in workmanship and materials under normal use for a period of Five Years from the date listed on the
- For repair under this warranty, our authorized service agent must be advised of the fault with the period of the warranty. This warranty covers parts and labor only under normal operations. Any defect resulting from natural causes (e.g., flood, hurricane, etc.) is not within this guarantee. This guarantee does not cover damage incurred by use of the unit not in accordance with the instructions, accidental damage, or being tampered with or serviced by unauthorized service agents.
- Monitor subjected to misuse, abuse, and neglect of this manual's content, non-instructional purposes, unauthorized repair or modifications will be excluded from this warranty.

The device requires no calibration.

The device is not repairable and contains no user serviceable parts.

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EMC Declaration

Table 2

Immunity Test	EC 60601-1-2 Test Level	Compliance Level
Electrostatic discharge (ESD) IEC 61000-4-2	±8 kV contact ±2 kV,±4 kV,±8 kV, ± 15 kV air	±8 kV contact ±2 kV,±4 kV,±8 kV, ±15 kV air
Electrical fast transient/burst IEC 61000-4-4 Surgel EC 61000-4-5	Power supply lines: ±2 kV 100 kHz repetition frequency line(s) to line(s): +0.5kV +1 kV.	Power supply lines: ±2 kV 100 kHz repetition frequency line(s) to line(s): +0.5kV +1 kV.
Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	0% 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% 1 cycle 70% 25/30 cycles Single phase: at 0 0% 250 cycle (50Hz)	0% 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0% 1 cycle And 70% 25/30 cycles Single phase: at 0 0% 250 cycle (50Hz
Power frequency magnetic field IEC 61000-4-8	30 A/m 50Hz/60Hz	30 A/m 50Hz/60Hz
Conduced RF IEC61000-4-6	150KHz to 80MHZ: 3Vrms 6Vrms (in ISM and amateur radio bands) 80% Am at 1kHz	150KHz to 80MHZ: 3Vrms 6Vrms (in ISM and amateur radio bands) 80% Am at 1kHz
Radiated RF IEC61000-4-3	10 V/m 80 MHz-2.7 GHz 80 % AM at 1 kHz	10 V/m 80 MHz-2.7 GHz 80 % AM at 1 kHz

Guidance and manufacturer's declaration - electromagnetic immunity

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tell| healTh

Arm Blood Pressure Monitor

Shenzhen Urion Technology Co.,Ltd.
Floor 4-6th of Building D, Jiale Science&Technology Industrial
Zone, No.3, ChuangWei Road, Heshulkou Community, MaTian
Street, GuangMing New District, 518106 Shenzhen, PEOPLE'S REPUBLIC OF CHINA

Manufactured for:Telli Health LLC 66 West Flagler Street - Suite 900, Miami,FL,33130 USA



REV.02

About blood pressure

Signal Strength Indicator:

H means signal is strong. L means signal is weak. means there is no signal. These marks will appear after finishing measurement and start to unload data









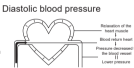
About blood pressure

Blood pressure is the pressure exerted the arteries

The systolic blood pressure value represents the blood pressured produced by contraction of the heart muscle.

The diastolic blood pressure value represents the blood pressured produced by relaxation of the heart muscle





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Care and Maintenance

Care for the main unit and blood pressure monitor cuff

 Keep the unit in the storage case when not in use. Clean the unit with soft dry cloth.

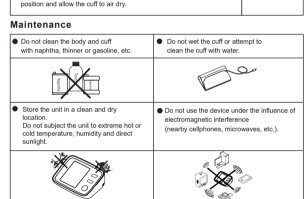
Do not use any abrasive or volatile Never immerse the unit or any

component in water Make sure the monitor is off prior to cleaning, a mixture of distilled water and 10 percent bleach could be used.

Using a spray bottle, moisten a soft cloth towel with the bleach or

detergent mix until it is fully saturated. Squeeze any excess moisture from the cloth to avoid any dripping or potential oversaturation of the Wipe all surfaces of the blood pressure monitor cuff thoro making sure to clean the inside and outside of the cuff. Be cautious

not to get any moisture in the main unit. Using a dry cloth, gently wipe away any excess moisture that may remain on the blood pressure cuff. Lay the cuff flat in an unrolled position and allow the cuff to air dry.



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EMC Declaration

IEC 60601-1-2: 2014 ME EQUIPMENT and ME SYSTEMS identification, marking and documents for Class B product. Instructions for Use:

The ME EQUIPMENT or ME SYSTEM is suitable for home healthcare environments and so on.

Warning: Don't use near active HF surgical equipment and the RF shielded room of an ME 329mm.n for magnetic resonance imaging, where the intensity of EM disturbances is high.

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 Warning: Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation. 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 the blood pressure monitor, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result. If any: A list of all cables and maximum lengths of cables (if applicable), transducers and other ACCESSORIES that are replaceable by the RESPONSIBLE ORGANIZATION and that are likely to affect compliance of the ME EQUIPMENT or ME SYSTEM with the requirements of Clause 7 (EMISSIONS) and Clause 8 (IMMUNITY). ACCESSORIES may be specified either generically (e.g., shielded cable, load impedance) or specifically (e.g., by MANUFACTURER and $\,$ EQUIPMENT OR TYPE REFERENCE).

If any: The performance of the ME EQUIPMENT or ME SYSTEM that was determined to be ESSENTIAL PERFORMANCE and a description of what the OPERATOR can expect if the ESSENTIAL PERFORMANCE is lost or degraded due to EM DISTURBANCES (the defined term "ESSENTIAL PERFORMANCE" need not be used).

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EMC Declaration

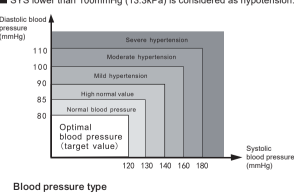
Table 3

Guidan	ice and n	nanufactur	er's declaration	- electroma			unity
Radiated RF IEC61000- 4-3 (Test	Test Freque ncy (MHz)	Band (MHz)	Service	Modulation	Modul ation (W)	Dist ance (m)	IMMUN TY TEST LEVEL (V/m)
specifications for ENCLOSU RE PORT IMMUNITY to RF wireless communications equipment)	385	380-390	TETRA 400	Pulse modulation 18 Hz	1,8	0.3	27
	450	430-470	GMRS 460, FRS 460	FM ± 5 kHz deviation 1 kHz sine	2	0.3	28
	710		LTE Band 13, 17 GSM 800/900,	Pulse modulation 217 Hz	0,2	0.3	9
	745	704 – 787					
	780						
	810		TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse modulation 18 Hz	2	0.3	28
	870	800 - 960					
	930						
	1720	1700 – 1990	GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse modulation 217 Hz	2	0.3	28
	1845						
	1970						
	2450	2400 – 2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation 217 Hz	2	0.3	28
	5240		WLAN 802.11 a/n	Pulse modulation 217 Hz	0,2	0.3	9
	5500	5100 – 5800					
	5785	. 5500	3/11				

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About blood pressure

- According to the blood pressure classification by the WHO/ISH,
- SYS lower than 100mmHg (13.3kPa) is considered as hypotension.



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Specification

Description	Automatic upper arm blood pressure monitor		
Display	LCD digital display		
Measuring principle	Oscillometric method		
Measuring localization	Upper arm		
Measurement	Pressure	0~299mmHg	
range	Pulse	40-199 pulses/min	
Accuracy	Pressure	3mmHg	
7.10041.409	Pulse	5% of reading	
	Pressure	3 digits display of mmHg	
LCD indication	Pulse	3 digits display	
indication	Symbol	Memory/Heartbeat/Low battery	
Memory function	2x90 sets memory of measurement values		
Power source	1pc 3.7V 1000mAh rechargeable lithium battery		
Automatic power off	in 3 minutes		
Main unit weight	Approx.219g(batteries not included)		
Main unit size	132mmx100mm×45mm		
Main unit lifetime	10,000 times under normal use		
Battery life	Could be used for about 70 times when charge is full		
Accessories	Cuff, instruction manual, lithium battery, USB cable,Wall charger		
0	Temperature	5°C~40°C	
Operating environment	Humidity	15%~93%RH	
	Air pressure	86kPa – 106kPa	
Storage environment	Air pressure: 86kPa ~106kPa; Temperature: -20°C ~ 55°C; Humidity:10% ~ 93%RH; avoid crash, direct sun or rain during transportation		
Expected service life	Five years		
Software version	UA1.0		

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EMC Declaration

Technical Description

- 1. All necessary instructions for maintaining BASIC SAFETY and ESSENTIAL PERFORMANCE with regard to electromagnetic disturbances for the excepted service life
- 2. Guidance and manufacturer's declaration electromagnetic emissions and Immunity.

Guidance and manufacturer's of emissions	declaration – electromagr
Emissions Test	Compliance
RF emissions CISPR 11	Group 1
RF emissions CISPR 11	Class B
Harmonic emissions IEC 61000-3-2	Class A
Voltage fluctuations / flicker emissions IEC 61000-3-3	Compliance

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FCC Statement

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.

Any changes or modifications not expressly approved by the party responsible for compliance could void the user's authority to operate the equipment.

Note: 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 reasonable protection against harmful interference in a residential installation. This equipmen 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 or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures: -Reorient or relocate the receiving antenna.

-Consult the dealer or an experienced radio/TV technician for help. FCC RF Exposure Information and Statement

which the receiver is connected.

When carrying the product or using it while worn on your body, either use an approved accessory such as a holster or otherwise maintain a distance of 5 mm from the body to ensure compliance with RF exposure requirements. Use of other accessories may not ensure compliance with FCC RF exposure guidelines.

-Increase the separation between the equipment and receiver. -Connect the equipment into an outlet on a circuit different from that to