

# SPIROMETER SP-10 BLUETOOTH

# Use and maintenance book

ATTENTION: Operators must read and understand this manual completely before using the product.



# Dear users, thank you very much for purchasing the SPIROMETER

Please read the User Manual carefully before using this product. The User Manual which describes the operating procedures should be followed strictly. Failure to follow the User Manual may cause measuring abnormality, equipment damage and human injury. The manufacturer is NOT responsible for the safety, reliability and performance issues and any monitoring abnormality, human injury and equipment damage due to users' negligence of the operation instructions. The manufacturer's warranty service does not cover such faults.

Owing to the forthcoming renovation, the specific products you received may not be totally in accordance with the description of this User Manual. We would sincerely regret for that

This product is medical device, which can be used repeatedly

### WARNING

For accuracy, it is recommended that the SPIROMETER should not be tested on the same testee for more than 5 times.

- The testee should breathe out all air during testing, don't exchange air or cough.
- Oon't use the device in environment with lower temperature
- Automatic nower off when there is no operation in one minute
- Please refer to the correlative literature about the clinical restrictions and caution
- This device is not intended for treatment
- Our company reserves the final elucidative right.

- 1.1 Instructions for Safe Operations
- Check the main unit and all accessories periodically to make sure that there is no visible damage that may affect patient's safety and monitoring performance. It is recommended that the device should be inspected weekly at least. When there is obvious damage, stop using the device.
- ♦ All maintenance must be performed by gualified service engineers ONLY. Users are not permitted to maintain it by themselves
- ♦ The SPIROMETER cannot be used together with devices not specified in User Manual. Only the accessory that is appointed or recommendatory by manufacture can be used with this device.

This product has been calibrated before leaving factory.

## 1.2 Warning

- Explosive hazard—DO NOT use the SPIROMETER in the environment with tinder such as anesthetic
- Please check the packing before use to make sure the device and accessories are totally in accordance with the packing list, or else the device may have the possibility of working abnormally.
- Don't use the device in environment with strong electromagnetic interference, direct breeze source, cold source and hot source
- Portable or mobile RF equipment with strong electromagnetic interference may influence the accuracy of this device.
- Improper disposal of device and its accessories and packing (include mouthpiece, plastic bags, foams and paper boxes) may cause environment pollution, please follow the local laws and regulations
- Please choose the accessories which are appointed or recommended by the manufacturer for avoiding device damage.
- Don't use the device with the turbine of the same kind product.
- DO NOT use the device when it is under charging state.
- The red and green indicators are all highlight in charging state, the red indicator goes out when the charge has finished. 1.3 Attention
- A Keep the SPIROMETER away from dust, vibration, corrosive substances, tinder, high temperature and moisture
- Generation → Generation → Generation
- A When it is carried from cold environment to warm or humid environment, please do not use it immediately.
- A DO NOT operate button on front panel with sharp things.
- A High temperature or high pressure steam disinfection to the device is not permitted. Refer to User Manual in the relative chapter (7.1) for cleaning and disinfection
- 😞 Do not have the SPIROMETER immerged in liquid. When it needs cleaning, please wipe its surface with medical alcohol by soft material. Do not spray any liquid on the device directly.
- a  $\,$  When cleaning the device with water, the temperature should be lower than 60°C.
- lacktrian End and the end and
- eta When data can't be displayed at all times or other cases happened during testing, press "repeated measure" key to remeasure or nower off to restart
- A The device has normal life for three years since the first electrified use
- When the data goes beyond the limits, the main screen shows "Error!".
- label{eq:2.1} & The device doesn't suit all users, if you can't get good measurement data, please stop using it.
- A The device needs to be calibrated once per year or less.
- A The device is forced SPIROMETER, according to the User Manual to use right to gain best result.

# 1.4 Contraindication

- 1.4.1 Absolute contraindication
- A The one with MI or shock in recent 3 months;
- A The one with serious cardiac function unstable or angina pectoris in recent 4 weeks;

- A The one with massive hemoptysis in recent 4 weeks:
- A The one who needs medication in epileptic seizure:
- A The one with uncontrolled hypertensive disease (SYS>200mmHg, DIA>100mmHg):
- A The one with aortic aneurysm
- A The one with serious hyperthyroidism
- 1.4.2 Relative contraindication
- A Heart rate >120 beats/min:
- A The one with pneumothorax or giant pulmonary bulla and not plan for surgical treatment:
- A The one with pregnancy:
- A The one with tympanic membrane perforation (need to block the ear canal of affected side before taking measurement)

5.1 View of the Front Pane

5.2 Assembly and disassemble

5.3 Accessories

1) A User Manual

2) A USB data line

3) A mouthpiece

5) PC software

6.1 How to use

6.1.2 Measu

6.1.1 Power on/off

6.1.3 Main interface

4) A power adapter

6) A nose clin (ontional)

Chapter 6 Operating Guide

parameter) interface as shown in Fig.4.

Case No.

Health status indicator -

"Denote value" under "Date managemen

005 12:23

100%

5.55

2.81

4.19

Figure 5 Other parameter interface

.....

6.1.4 Menu

FEV1%

FEF25

FEF75

FEF2575

the bottom, counterclockwise rotate to lock it

2)Turbine disassembly: clockwise rotate the turbine, gently pull it out

3)Mouthpiece assembly: insert the mouthpiece into the turbine port directly

must meet the requirements of EN60601 related standards and have the CE mark.

(1) After assembly, long press "power on" key to turn on the device.

personal information, after exit, it will return to (Testing) interface.)

minimal amount of time, wait for a few seconds, the device will enter (Mair

Do you want to edit personal information ?

Figure 2 Selective interface

FVC

EEV1

battery status, time, case number and health status indicator, as shown in Fig.4.

005 12:23

3 50

3.15

Figure 4 Main parameter interface

health status, correct settings of personal information is the key to obtain accuracy ratio. Besides, this interface can also display

image, i.e. Compare the measured value with the reference value in same situation. When the value is lower than 50%, only red

indicator is displayed, which means testee should pay attention and go to hospital in time. When the value is in range from 50%-80%

red and yellow indicator are displayed, which means it should be noticed. When the value is higher than 80%, all red, yellow and

green indicator are displayed, which means healthy. The determinate item of health status indicator is optional, it can be set in

V-L

Figure 6 Flow rate-volume chart

functions such as modify personal information, data management, device setting, power off can be realized. Press "Up" or "Down"

key to move the selection toolbar to the item that need to modified, then press "Confirm" key to enter the sub-menu. See the

Under (Testing) or (Main interface), press confirm key to enter (Menu) interface as shown in Fig.8. Under the interface,

d.Under (Main parameter) interface, press "Up" or "Down" key will enter (Other parameter) (Flow rate-volume chart)

c.Other parameter interface: display four parameters except the main parameter, as shown in Fig.5.

(Volume-time chart) in turn, as shown in Fig.5, 6, 7, The four interfaces above are (Main interface)

a.Main parameter interface: display the ratio of predicted value and measured value of three main parameters. Ratio reflects

**b.Health status indicator:** indicate the ratio of measured value and the predicted value, display the testee health condition in

(2) When device is powered on, long press "power off" key to turn it off.

Charging in

1

Figure 1 Front panel view

1)Turbine assembly: Hold the turbine, align the arrowhead of the turbine with the triangular shape on the shell, gently insert it to

△ Other type adapter should meet the following conditions: output voltage:DC 5V; output current≥500mA, the power adapter

(1) The device is in (Selective interface) after turn on as shown in Fig.2, press "up" or "down" key to select "No", then press

"confirm", key to enter (Testing) interface as shown in Fig.3. (Note: If select "Yes", it will enter (Personal information) interface to edit

(2) In (Testing) interface, breath in fully, seal the lips around the mouthpiece and blast all air out as forcefully as possible in a

Testina.

Figure 3 Testing

Ratio of me

predicted value

Predicted value is a reference under the

e been set. It is a popularize value

uation that values (gender age, height, etc.)

red value and

L-T

Figure 7 Volume-time char

- The one with RTI recently (less than 4 weeks):
- $\Theta$  The one with hypoimmunity.

Patients of respiratory communicable disease or infectious disease shall not take lung function examination in the acute stage. The one with low immunity is not appropriate to take the examination also. If it is necessary, disease control and protection shall be strictly followed

### 1 5 FMC declaration

- A When this device is installed or putted into service, EMC should be paid more attention, as the portable and mobile RF communications equipment with higher EM interference can affect this device.
- A The internal components and cables should not be changed, as this may decreased IMMUNITY of the device.
- A The SPIROMETER should not be used adjacent to or stacked with other equipments.

Forced Vital Capacity is the maximum expiration after taking a full breath, it's an important examination content in chest-lung disease and respiratory health, and it is indispensable testing project in modern Pulmonary inspection. At the same time, it has great significance in respiratory diseases, differential diagnosis, treatment evaluation and selection of surgical indications. Thus, with the rapid development of clinical respiratory physiology, clinical applications of lung capacity inspection are also gaining popularity. The SPIROMETER is small in volume, low in power consumption, convenient in operation and portable. With high-definition display screen, the device is concise and fashion. It is only necessary for patient to breath in fully and seal the lins around the mouthpiece and blast the air out in best times for measure, then the display screen will directly show the Forced Vital Capacity (FVC). Forced Expired Volume in one second (FEV1), Peak Expiratory Flow (PEF) with the high veracity and repetition.

### 2.1 Features

- 1) Ultra-thin design, concise and fashion.
- 2) Small in volume, light in weight and convenient in carrying.
- 3) Low power consumption
- 4) TFT display.
- 5) Reflect lung function by measuring FVC, FEV1, PEF etc.
- 6) Take the function of wireless trans

### 2.2 Major Applications and Scope

The SPIROMETER is a hand-held equipment for examining lung function. The product is fit for hospital, clinique, family for ordinary test. It's only required that the user operates it according to user manual, no need for specialized training so the operation of the device would be as simple and easy as possible.

### 2.3 Environment Requirements

- Storage Environment:
- Temperature: -40°C~+55°C
- Relative humidity: ≤95%
- Atmospheric pressure: 500hPa~1060hPa
- Operating Environmen
- Temperature: +10°C~+40°C
- Relative Humidity: ≤80%

# Atmospheric pressure: 700hPa~1060hPa

Chapter 3 Principle

Firstly, testee deep inspires, then seals the lips around the mouthpiece and blasts all air out as forcefully as possible, the exhalant gas transforms to rotary airflow by turbine, then makes the blade rotate. The reception part of the infrared pair diodes (one is for infrared emission, the other is reception) towards to the blade is used for receiving the infrared ray, when the blade rotates, the received ray strength of the reception diode will be different as the difference of the blade angle, so form the various signal of same proportion in reception diode, which forms acquisition signal by SCM after processing. At last, various parameters to be measured formed from the information which were processed by the microprocessor, and displayed from the screen.

- 4.1 Main Performance
- Forced Vital Capacity (FVC), Forced Expired Volume in one second (FEV1), the ratio of FEV1 and FVC (FEV1%), Peak expiratory flow (PEF), 25% flow of the FVC (FEF25), 75% flow of the FVC (FEF75) and average flow between 25% and 75% of the FVC (FEF2575) can be measured. Besides, the testee condition can be shown by the ratio of the measured value and the predicted value
- Flow rate-volume chart, volume-time chart display.
- Data memory, delete, upload and review.

Volume accuracy: ±3% or 0.05L (whichever is greater)

Power supply: DC3.7V 820mAh rechargeable lithium battery

According to the MDD 93/42, the classification of this medical device: II a.

The degree of protection against electroshock: Type BF applied part.

The type of protection against electroshock: Internally powered equ

Flow accuracy: ±5% or 0.2L/s (whichever is greater)

- Trend chart display.
- Calibration

4.2 Main Parameters

Volume Range: 10L

Flow range: 0 L/s~16 L/s

Working current: 60mA

Classification

EMC: Group I Class B.

International Protection: IP22.

- Information prompts when volume or flow goes beyond the limits.
- Automatic power off when there is no operation in one minute.
- Rechargeable lithium battery and with charging tips. Battery power display.

following steps for details:

Menu
Personal Information
Data Management
Settings
Power Off
Exit

Figure 8 Menu interface

Personal Information				
Number	36			
Gender	FEMALE			
Age	20			
Height / cm	160			
Weight / kg	50			
Nation	ERS			
Smoker	NO			

Figure 9 Personal information interfac

### a. Personal in

Under (Menu) interface, select "Personal information" to enter its interface as shown in Fig.9, in which user can edit patient information (Note: Under (Selective interface) as shown in Fig.2, if selected "Yes", you can enter (Personal information)interface also.).

### (1) Case number

"Number" is the case number displayed at present. For example, if you are the 36th testee, the "Number" will be 36, Case number can increase automatically, no need to set manually.

## (2) Gender setting

Under (Personal information) interface, press "Up" or "Down" key to move the selection toolbar to "Gender", then press "Confirm" key to select "female" or "male

### (3) Setting of age, height, weight

Under (Personal information), select "Age" to enter (Age edit) interface, as show in Fig.10. Press "Up" or "Down" key to change the value. At each pressing of "Up" or "Down" key, the value will plus or minus 1. When long press the "Up" or "Down" key, the value will increase or decrease continuously. Press "Confirm" key to back to (Personal information) interface.

The modification of "Height" and "Weight" is similar to the "Age". In which, range of "Age" is 6~100 years old, range of "Height" is 80~240 cm. range of "Weight" is 15~250 kg.



Figure 10 Age edit interface

### (4) Nation setting

The modification of "Nation" is similar to the "Gender". The standard of predicted value can be set under "Nation" interface. which including ERS. KNUDSON and USA. ERS is the European standard, KNUDSON is the Asian standard, USA is the American ctandard

### (5) Setting of smoker and drug

The modification of "Smoker" and "Drug" is similar to the "Gender", in which patient information of smoker and drug can be modified

For the display of screen is limited, the device won't display all items at the same time. When selection toolbar moved to "Smoker", press "Down" key, the item of "Drug" and "Exit" will appear, as shown in Fig. 11, 12.

### (6) Exit

Under (Personal information) interface, select "Exit" to return to (Menu) interface.



Figure 11

### b.Data managemer

Under (Menu) interface, select "Data management" to enter (Data management) interface, as shown in Fig.13. Under the interface, functions such as review, view trend curve, delete data, denote value setting can be realized



Figure 13 Data management interface



Figure 12

160 50

ERS

### (1) Review function

Under (Data management) interface, select "Review function" to enter (Case selection) interface as shown in Fig.14, press "Up" or "Down" key (long press is available) to change case number, then press "Confirm" key, the device will enter (Main interface) and display history data on it. Under (Main interface), press "Up" or "Down" key continuously can review data in adjacent case number, press "Confirm" key to return to (Menu) interface.

# (2) Trend curve

Under (Data management) interface, select "Trend Curve" to enter (Trend curve selection) interface as shown in Fig.15. Select the determinant parameter, then press "Confirm" key to enter (Trend curve display) as shown in Fig.15. The curve is a summary of stored data for selected parameter. It displays the change trend in form of visual image, which is convenient for comparison, If the data is too much, press "Up" or "Down" key to browse all data trend curves orderly. Press "Confirm" key to return to (Data management) interface.

Please select determinant of trend curve
FVC
FEV1
PEF
FEV25
FEV75
FEV2575



Figure 15 Trend curve selection interface

Figure 16 Trend curve display interface

# (3)Delete data

Under (Data management) interface, select "Delete data" to enter (Delete data) interface as shown in Fig.17. If choose "Yes",

the screen displays "waiting...", all data will be deleted, then return to (Data management) interface. If choose "No", it will return to (Data management) interface directly.

Are you sure to delete all the data ?

Figure 17 Delete data interface

Which one to decide the

Select one parameter to decide the denote value, after that, it will automatically return to (Data management) interface.

FE\/1%

Under (Data management) interface, select "Denote value" to enter (Denote value setting) interface as shown in Fig.18.

full, it will disolay (Memory full) interface as shown in Fig.27. If you select "Yes", it will enter (Delete data) interface: if you select "No". Under (Data management), select "Exit" to return to (Menu) interface. it will enter (Menu) interface. Settings Language

Figure 20 Language setting interface

8

Figure 22 Minute setting interface

Calibrate

Figure 24 Calibrate interface

language, Bluetooth on/off, time and calibration, and view device information can be realized.

Under (Settings) interface, select "Language" to enter (Language setting) interface as shown in Fig.20, Select "English", the device language will be English, select "中文", the device language will be Chinese, after selected, it will automatically return to

Move selection toolbar to "Bluetooth", press "Confirm" key to select "ON" or "OFF" that can turn on or off the Bluetooth

Under (Settings) interface, select "Time" to enter (Time setting) interface as shown in Fig.21, Select "Minute" to enter

The operation of "Hour", "Day", "Month", "Year" is similar to the "Minute". The "Week" will be calculated according to "Year",

Under (Settings) interface, select "Calibration" to enter (Calibration setting) interface as shown in Fig.23. Select 2L or 3L

Under (Calibrate) interface, push the syringe once, the device will display "REPEAT", then push the syringe once again. After

twice correct continuous operation, the calibrating will be succeed, and the device will display "OK!". Finally the interface will

jump to the former interface before calibration (The former interface: If the device is calibrated after measurement completed, it

until succeeded. If the device displays "Select right volume", please confirm whether the volume of syringe and calibration

selection is accordant, then repeat the calibrating until succeeded. If you need to stop calibrating, just press the "Confirm" key to

Under (Calibration setting) interface, select "Adjust" to enter (Adjusting) interface, as shown in Fig.25. Press "Up" or "Down"

key to change the value (long pressing is available), then press "Confirm" key to return to (Adjusting confirm) interface, as shown

in Fig.26. Selecting "Yes" will save adjusted value, selecting "No" will cancel the setting, then the device will return to (Calibration

🗥 Note: The value determines the accuracy of measurement, please do NOT change it randomly. After the turbine has been

replaced, calibration shall be applied for inputting parameters of new turbine, which guarantees the accuracy of measurement

If the device displays "Error! Please repeat", it indicates something wrong with the operation, please repeat the calibrating

will return to (Settings) interface; if calibrated before measurement completed, it will return to (Testing) interface.).

(Minute setting) interface, as shown in Fig.22. Press "Up" or "Down" key to change the value (long pressing is available), then

"Month" and "Day", which does not need to set manually. Then select "Exit" to return to (Settings) interface

Time Setting

Figure 21 Time setting interface

based on the volume of syringe, then enter to (Calibrate) interface as shown in Fig.24.

Calibration

Figure 23 Calibration setting interface

Figure 18 Denote value setting interface

Under (Menu) interface, select "Settings" to enter (Settings) interface as shown in Fig.19. Under this interface, settings of

(5)Exit

Figure 19 Setting interface

module (If there is no Bluetooth module in the device, the operation is invalid).

press "Confirm" key to return to (Time setting) interface.

Day Ionth

′ear

Adius

exit to the former interface before calibration.

setting) interface.

after turbine replaced.

(4)Denote value

(1) Language setting

(Settings) interface.

(2) Bluetooth

(3) Time setting

(4) Calibratio

### Table 1 Guidance and manufacturer's declaration – electromagnetic emission Emission tes RE omi Group 1 CISPR 11 RF emission Class B CISPR 11 Harmonic emissions Class A FC 61000-3-2 Voltage fluctuations/ flicker emissions Comply EC 61000-3-3

The malfunction of the device

The battery is not full charged.

drained away.

The battery is broken.

The battery is broken.

Follow instructions for use

overing Protection rate

VEEF disposal

Full-powe

Low-power

vpe BF Applied part.

tatus indicator bar

midity limitat

nperature limit

ragile, handle with care

Keep in a cool, dry place

his way up

Date of manufactur

1anufacturer

erial number

harging indicator

ledical device

Product code

ot number

urn the turbine clockwise to unloci

urn the turbine counterclockwise to lock

uthorized representative in the European community

Description

Forced vital capacity

Peak expiratory flow

25% flow of the FVC

75% flow of the FVC

manufacturer. Otherwise, degradation of the performance of this equipment could result.

The ME EQUIPMENT or ME SYSTEM is suitable for home healthcare environment

imaging, where the intensity of EM disturbances is high.

they are operating normally.

disturbances for the excepted service life

Instructions for use

FEV1/FVC×100

Forced Expired Volume in one second

Average flow between 25% and 75% of the FVC

Warning: Don't near active HF surgical equipment and the RF shielded room of an ME system for magnetic resonance

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

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 equipment, including cables specified by the

all necessary instructions for maintaining BASIC SAFETY and ESSENTIAL PERFORMANCE with regard to electromagnetic

٩

portated by

Unit

1/s

L/s

L/s

mospheric pressure limit

The display disappears sudden

The device can not be used for

The battery can not be full

charged even after 10 hours

The device has built-in wireless

nodule, but can't achieve

Chapter 9 Key of Symbols

6

CE

IP22

A

R

Ŕ

Error

0.0

(2)

m

----

SN

-8-

EC REP

MD

REF

LOT

Chapter 10 Parameter I

Measured parameters

Parameter

FVC

FFV1

PEF

FEV1%

EFE25

FEF2575

FEF75

Appendix I

2

full time after charge.

charging time.

reless trai

Are you sure to save

Figure 26 Adjusting confirm interface

145

Figure 25 Adjusting interface

Under (Settings) interface, select "About" to enter (About) interface. User can view device name and software version. Press

Under (Menu) interface, select "Exit" to return to (Main interface). If the measurement is not completed before enter

Measurement of the device is repeatable. Long press "Repeated measure" key to enter (Testing) interface. When the memory is

ny io full

Figure 27

A For device charging, connect it with the power where easy to be cut off, after charging completed, unplug the power adapter

...

Figure 28

2) Press the corresponding key to achieve upload data, delete case, print information, background, select language, switch PDF

A Excessive ambient light may affect measurement accuracy. It includes fluorescent lamp, dual ruby light, infrared heater, direct

Using medical alcohol to wipe the device for disinfecting, nature dry or clean it with clean soft cloth. It's necessary to clean the

turbine periodically for accuracy, keep the diaphaneity of the lucency part, and keep it away sundries (such as hair or lesser sediment). Immerse the turbine in disinfectant after use, clean it with clean water and dry standing vertically after soaked a few

minutes (but don't make the turbine rinsed with water directly), this type doesn't bring pollution to environment. (Note: The

3) Recharge the battery soon after the over-discharge. The device should be recharged every six months when it is not regular

4) The device needs to be calibrated once a year (or according to the calibrating program of hospital). It also can be performed at

1) The packed device can be transported by ordinary conveyance or according to transport contract. The device can not be

The packed device should be stored in room with no corrosive gases and good ventilation. Temperature: -40°C~+55°C: Relative

Solution

emeasure according to the user manual.

Press "Repeated Measure" key to remeasur

Delete the current case and remeasure.

Operate normally according to the use

Please contact the local service center.

or power off to restart.

Please charge the battery

anual.

Possible Reason

The start speed is too low, the device doe

The malfunction of the device

The malfunction of the device

The power turned off abnormally.

used. It can extend the battery life following this guidance. If the battery is broken, DO NOT try to maintain it by yourself,

1) Connect the device with computer by data line — then the device should be under charging state.

Install the PC software in the computer, then the following figure will appear after completing

2) Connect the device with power supply by power adapter, then the device should be under charging state.

1) Connect the device with computer by data line double press the icon to open the PC software procedure

3) Press "Exit" to exit the software, unplug the data line from the computer to achieve uploading.

A Intense activity of the subject or extreme electrosurgical interference may also affect the accuracy.

Please clean and disinfect the device after using according to the User Manual (7.1).

Please clean and disinfect the device before using according to the User Manual (7.1).

ot measure.

peration is wrong.

2) Please recharge the battery when the screen shows low-power (the battery power is

B Please check the device before using, and confirm that it can work normally.

 $\beta$  It is recommended that the device should be measured in room.

Under (Calibration setting) interface, select "Exit" to return to (Settings) interface.

Under (Menu) interface, select "Power off", the device will shut down

Note: If there is no operation within 1 minute, the device will power off automatically.

Under (Settings) interface, select "Exit" to return to (Menu) interface

(5) About device

(6) Exit

e.Exit

6.1.6 Charge

to cut off from nower

6.1.7 Upload Data

6.2 Attention

A Rechargeable lithium battery.

sunlight and etc.

7.1 Cleaning and Disinfection

disinfectant is 75% alcohol).

7.3 Transportation and Storage

Humidity: ≤95%.

Trouble

neasurement for a long time

and the data can't be displayed

Chapter 8 Troubleshoot

The device can't finish

The figure is wrong and

norderly.

7.2 Maintenance

There are two kinds of charging methods:

format, set the testee information etc.

Chapter 7 Maintenance, Transportation and Stor

please contact us or the local service center.

the state-appointed agent or just contact us for calibration.

transported mixed with toxic, harmful, corrosive material.

The device can not be powered Low battery or no pow

d Power off

6.1.5 Repeated measure

"Confirm" key to return to (Settings) interface.

(Main interface), it will return to (Testing) interface.

The device is set to automatic power off

The battery is drained away or almost

The wireless module is broken, or the

ansmission route has problem

ledical Device compliant with Directive 93/42/EEC

aution: read instructions (warnings) carefully

Measured value goes beyond the limits

hen there is no operation in one minute

Please contact the local service center.

Please contact the local service center.

lease contact the local service center

ase contact the local service center

ease charge the battery.

Please recharge the battery.

nrma

Meaning

Guidar	nce and manufacturer's declaration – electro	omagnetic immunity			
Immunity test	IEC 60601-1-2 test level	Compliance level			
Electrostatic discharge (ESD)	±8 kV contact	±8 kV contact			
IEC 61000-4-2	±2 kV, ±4 kV, ±8 kV, ±15 kV air	±2 kV, ±4 kV, ±8 kV, ±15 kV air			
Electrical fast transient/burst	$\pm 2$ kV for power supply lines	$\pm 2$ kV for power supply lines			
IEC 61000-4-4	$\pm$ 1 kV signal input/output	Not applicable			
	100 kHz repetition frequency	100 kHz repetition frequency			
Surge	$\pm$ 0.5 kV, $\pm$ 1 kV differential mode	$\pm$ 0.5 kV, $\pm$ 1 kV differential mode			
IEC 61000-4-5	$\pm$ 0.5 kV, $\pm$ 1 kV, $\pm$ 2 kV common mode	Not applicable			
Voltage dips, short interruptions	0 % UT; 0,5 cycle. At 0°, 45°, 90°, 135°,	0 % UT; 0,5 cycle. At 0°, 45°, 90°, 135°, 180°,			
and voltage variations on power	180°, 225°, 270° and 315°.	225°, 270° and 315°.			
supply input lines	0 % UT; 1 cycle and 70 % UT; 25/30	0 % UT; 1 cycle and 70 % UT; 25/30 cycles;			
IEC 61000-4-11	cycles; Single phase: at 0°.	Single phase: at 0°.			
	0 % UT; 250/300 cycle	0 % UT; 250/300 cycle			
Power frequency magnetic field	30 A/m	30 A/m			
IEC 61000-4-8	50Hz/60Hz	50Hz/60Hz			
Conducted RF	3 V	3 V			
IEC61000-4-6	0,15 MHz - 80 MHz	0,15 MHz - 80 MHz			
	6 V in ISM and amateur radio bands	6 V in ISM and amateur radio bands between			
	between 0,15 MHz and 80 MHz	0,15 MHz and 80 MHz			
	80 % AM at 1 kHz	80 % AM at 1 kHz			
Radiated RF	10 V/m	10 V/m			
IEC61000-4-3	80 MHz – 2,7 GHz	80 MHz – 2,7 GHz			
	80 % AM at 1 kHz	80 % AM at 1 kHz			

Table 3

	Guidance ar	nd manufactur	er's declaration –	electromagnetic	immunity	
	Test Frequency (MHz) 385	Band (MHz) 380 – 390	Service	Modulation	IEC 60601-1-2 Test Level (V/m) 27	Compliance level (V/m) 27
	385	380-390	TETRA 400	modulation 18 Hz	27	27
Radiated RF IEC61000-4-3 (Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipment)	450	430 –470	GMRS 460, FRS 460	FM ±5kHz deviation 1 kHz sine	28	28
	710 745 780	704 – 787	LTE Band 13, 17	Pulse modulation 217 Hz	9	9
	810 870 930	800 – 960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse modulation 18 Hz	28	28
	1720 1845 1970	1700 -1990	GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse modulation 217 Hz	28	28
	2450	2400 –2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse modulation 217 Hz	28	28
	5240 5500 5785	5100 - 5800	WLAN 802.11 a/n	Pulse modulation 217 Hz	9	9

Guidance and manufacturer's declaration - electromagnetic Immunity							
Radiated RF Test IEC61000-4-39 Frequency (Test specifications for		Test		IEC 60 Test L (A/m)	evel	Compliance level (A/m)	
ENCLOSURE PORT	30 kHz	CW		8		8	
IMMUNITY to proximity magnetic	134,2 kHz	Pulse n 2.1 kHz	nodulation	65		65	
fields)	13,56 kHz	Pulse n 50 kHz	nodulation	7,5		7,5	
Number	Model		Cable length	(m)	Mask or no	Remark	
1	Power adapter cable		1.50	(11)	YES	/	

Attention: With the exception of energy exchange and cables sold by manufacturers of lung function devices as spare parts for internal components, the use of accessories and cables other than those specified will result in increased product emission or reduced anti-interference.

The following cable types must be used to ensure compliance with interference radiation and immunity standards. Table: Cable overview



Disposal: The product must not be disposed of along with other domestic waste. The users must dispose of this quipment by bringing it to a specific recycling point for electric and electronic equipment.

### GIMA WARRANTY TERMS

The Gima 12-month standard B2B warranty applies