Preface

Please read the User Manual carefully before using this product. The operating procedures specified in this User Manual should be followed strictly. This manual describes in detail the operation steps which must be noted, the procedures which may result in abnormality, and possible damage to the product or users. Refer to following chapters for details. Failed to follow the User Manual may cause measuring abnormality, device damage or personal injury. The manufacturer is NOT responsible for the safety, reliability and performance issues of such results due to user's negligence of this user manual for using, maintenance or storage. The free service s and repairs do not cover such faults either.

The content in this user manual complies with real product. For software upgrade and some modifications, the content in this user manual is subject to change without prior notice, and we sincerely apologize for that.

Attentions

Before using this product, the safety and effectiveness described in the following shall be considered:

- Type of protection against electric shock: class I (AC power supply), internal powered equipment (power supplied by battery)
- Degree of protection against electric shock: type CF, defibrillation-proof function applied part
- Working mode: continuous running equipment
- Enclosure protection class: IPX0
- Measurement results shall be described by professional doctor combined with clinical symptoms.
- The using reliability depends on whether the operation guide and maintenance instructions in this user manual is followed.
- Service life:10 years
- Date of manufacture: see the label
- Contraindications: none

Warning: To ensure the device safety and effectiveness, please use the company recommended accessories. The maintenance and repair of the device should be done by

professional personal specified by the company. It is forbidden to refit the device.

Responsibility of the operator

- The device must be operated by a professionally trained medical staff, and kept by a special person.
- The operator should read the User Manual carefully before use, and strictly follow the operating procedure described in the User Manual.
- The safety requirements have been fully considered in product designing, but the operator can not ignore the observation of the patient and device.
- > The operator is responsible for providing the information of product use to the company.



Responsibility of the company

- > The company supplies qualified products to user in accordance with enterprise standard.
- > The company installs and debugs the equipment and trains the physicians by contract.
- > The company performs device repair in warranty period (a year) and maintenance service after warranty period.
- > The company responds timely to the user's request.

The user manual is written by our company. All rights reserved.

Statement

Our company owns all rights to this unpublished work and intends to maintain it as confidential information. This user manual is used only for reference of operation, maintenance, or repair of our device. No part of this can be disseminated to others. And our company takes no responsibilities for any consequences and liabilities caused by using this user manual for other purposes.

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Our company owns the final explanation right to this user manual, and reserves the right to change the content of this user manual without prior notice, and the rights to change product technology and specification.

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Chapter1 Overview

1.1 Overview

This product is a kind of electrocardiograph, which is able to sample 12-lead (E12), or 18-lead (E18M) ECG signals simultaneously and print out the ECG waveform with thermal printing system. Its functions are as follows: recording and displaying ECG waveform in auto/manual mode; measuring ECG waveform parameters automatically, and automatic analysis and diagnosis; pacing ECG detections; prompt for electrode-off and out of paper; optional interface languages(Chinese/English, etc.); built-in lithium battery, powered either by AC or DC; arbitrarily select the rhythm lead to conveniently observe abnormal heart rhythm; case database management, etc.

1.2 Intended use

The device is used in hospitals or professional medical institutions. Medical personnel with professional technical training can use this device to collect and record ECG signals of human body.

1.3 Main technical specifications

1.3.1 Environment conditions

Operation:

```
a). Environment temperature: 5°C~40°C
```

b). Relative humidity: 25%~95% (no condensation)

- c). Atmospheric pressure: 700 hPa~1060 hPa
- d). Power supply:

Voltage: 100-240 V \sim

Frequency: 50 Hz, 60 Hz

```
Input power: ≤150 VA
```

Battery: 14.8 V, 5200 mAh rechargeable lithium battery

Transportation and Storage:

- a). Environment temperature: -20 °C~+55 °C
- b). Relative humidity: ≤95%
- c). Atmospheric pressure: 500 hPa~1060 hPa
- 1.3.2 Input way: Floating and defibrillation protection
- 1.3.3 Lead: Standard 12 leads (E12)/ Standard 18 leads (E18M)
- 1.3.4 Patient leakage current: <10µA
- 1.3.5 Input impedance: $\geq 100M\Omega(10Hz)$

1.3.6 Frequency response:

Test	Rated input	Input frequency and waveform	Relative output		
	amplitude	1 1 2	response		
А	1.0	0.01Hz~0.67Hz, Sine wave	+ 5%, -30% ^a		
A	1.0	0.67Hz~40Hz, Sine wave	+ 5%, -10% ^a		
в	0.5	40Hz~100Hz, Sine wave	+ 5%, -10% ^a		
D	0.25	100Hz~150Hz, Sine wave	+ 5%, -20% ^a		
C 0.5		150 Hz ~ 300 Hz, Sine wave	+ 5%, -30% ^a		
0.5 350 Hz ~ 500 Hz, Sine wave		+ 10%, -100% ^a			
D	1.5	5 ≤ 1 Hz,200ms, Triangle wave +0 %, -10 % ^b			
^a relative to 10Hz ^b relative to 200 ms					

1.3.7 Time constant: ≥3.2s

1.3.8 CMRR: >123dB

>140 dB(Open filter)

1.3.9 Filter: AC Filter(50Hz/60Hz), EMG Filter(25Hz/35Hz (-3 dB)), DFT Filter, Low-pass Filter

1.3.10 Recording way: Thermal printing system

1.3.11 Specification of recording paper: 210 mm(W)×20 m(L) high-speed thermal paper

1.3.12 Time base selection(paper speed):

5mm/s、6.25mm/s、10mm/s、12.5 mm/s、25 mm/s、50 mm/s, error: ±5%

1.3.13 Gain control(sensitivity):

1.25, 2.5, 5, 10, 20, 40, 10/5, 20/10mm/mV and automatic, accuracy is $\pm 5\%$

Standard sensitivity: 10 mm/mV±0.2 mm/mV

1.3.14 Auto record: record setup according to auto record format and mode, automatically change leads, automatically measure and analyze.

1.3.15 Rhythm record: record setup according to rhythm record format and mode, automatically measure and analyze.

1.3.16 Manual record: record according to manual record format.

1.3.17 Measurement parameters: HR, PR Interval, P Duration, QRS Duration, T Duration, QT/ QTc Interval, P/QRS /T Axis, R(V5), S(V1), R(V5)+S(V1) amplitude

1.3.18 Product safety type: Class I type CF defibrillation-proof function applied part

1.3.19 Polarization resistance voltage: ± 950 mV, sensitivity variation range: $\pm 5\%$

1.3.20 Noise level: $\leq 12 \mu Vp-p$

1.3.21 ECG signal input sampling frequency: 32 kHz

1.3.22 Waveform data processing sampling frequency: 1 kHz

1.3.23 Sampling precision: 24 bit

1.3.24 The minimum detection signal: 10 Hz, 20 μ V(peak-peak value) deflected sinusoidal signal can be detected

1.3.25 Pacing detection channel: All channel

1.3.26 Pacing signal sampling frequency: 32kHz

1.3.27 Accuracy of input signal: The overall system error, ±5%.

1.3.28 Amplitude quantization: ≤5µV/LSB

1.3.29 Interchannel time deviation: <100 µs

1.3.30 Calibration voltage:1mV $\pm 2\%$

1.3.31 Input loop current:≤0.01uA

1.3.32 50Hz interference suppression filter: \geq 20dB.

1.3.33 Pace pulse display capability: ±2mV~±750mV, 0.1ms~2.0ms

1.3.34 Fuse specification: 2pcs φ5×20mm AC delay insurance: T3.15AH250V

1.3.35 Size: 340 mm(L)×320 mm(W)×91mm(H)

1.3.36 Net Weight: 5 kg

1.4 Main Characteristics

1.4.1 Display with 1280*800 dots 10.1 inch high resolution color LCD, operate either by touch screen or function buttons, which is convenient and quick.

1.4.2 Sync collection for 12-lead ECG(E12)/18-lead ECG(E18M), support for 12-lead(E12)/ 18-lead (E18M) and Cabrera-lead waveform display, adopt digital signal processing technology and get high-quality ECG via power frequency filter, baseline filter, EMG filter and Low-pass Filter for ECG signals.

1.4.3 Display of 3/6/12/18-lead (E18M) ECG waveform on one screen, and HR value, print mode, sensitivity, paper speed, filter state, clock, battery level, background gridlines, measured data and interpretation information, etc. Prompting function for lead-off and overload, system working state.

1.4.4 The device can be powered either by AC or DC(can adapt to 50/60Hz AC frequency), with built-in rechargeable lithium battery and charging circuit, perfect battery overcurrent and overvoltage protection circuit.

1.4.5 In optimal DC state, up to 10-hour standby time, continuous print more than 3-hour, record up to 1000 ECG waveform(commonly, it is 3s case), which meets the requirements of visiting a patient at home and body examination.

1.4.6 Built-in thermal printer, support for Auto M*N, Auto M*N+1, Auto M*N+2, Auto M*N+3, rhythm M line, manual and other printing modes and formats. The printed content contains time, paper speed, sensitivity, calibration signal, name of lead, filter state and patient's information. Information including printed waveform length, output measurement parameter, diagnostic conclusion, superposition QRS waveform, histogram, trend chart and interval list, can be set, and with time print function and auto arrhythmia print function, which meets different requirements.

1.4.7 With the functions of auto-measurement and auto-interpretation for routine ECG parameters, provide measurement results and auto-diagnosis conclusion for HR, PR Interval, P Duration, QRS Duration, T Duration, QT/QTc Interval, P/QRS/T Axis, R(V5), S(V1), R(V5)+S(V1) amplitude, Cornell Index, etc. which reduces the doctor's burden.

1.4.8 The built-in large-capacity memory can store at least 4000 medical records, making it easy for doctors to review medical records and statistical information.

1.4.9 Multi-language(Chinese, English) interface and report. Full touch screen with buttons operation, built-in virtual keyboard, support Chinese and English input methods.

1.4.10 With Wi-Fi wireless communication function.With functions of LAN, USB cable transmission. Automatically upload cases, download reports and print them.

1.4.11 Support external USB standard keyboard, mouse, scanner, printer.

1.4.12 Historical medical records can be reviewed, inquired, modified, transmitted, printed, lead correction, exported to other electronic file formats (dat, pdf, xml, bmp, jpeg, png etc.).

1.5 Software overview

Name of software: native embedded software

Software specification: none

Software version: V1.1.5

Version naming rules: V<major version number>.<minor version number>.<revision version number>

The version of the software can be obtained in "About".

Involved algorithm:

Name: ECG algorithm

Type: mature algorithm

Use: by processing and analyzing the static ECG data, measurement parameters such as HR of ECG and automatic interpretation items are obtained.

Clinical function: provide measurement parameters such as HR of ECG and automatic interpretation items to assist the physician in diagnosing cardiovascular disease. Automatic measurement parameters and interpretation results are for reference by the physician only, and can not be used as the sole basis for clinical diagnosis. The final diagnosis needs to be combined with the clinical symptoms.

Chapter2 Safety Precautions

2.1 Ensure that the device is placed on a flat level worktable. Avoid too strong vibration or impact when moving it.

2.2 When working with AC power, the power cord must be 3-core, the frequency and voltage value of the AC power source must match the identification on the manual and have sufficient capacity.

2.3 A perfect power system and grounding is need in room.

Mwarning: To avoid the risk of electric shock, this equipment must only be connected to

a power supply network with a protective ground.

2.4 If there are any questions for the integrality of protective grounding cable or the reliability of protective grounding cable connection can not be guaranteed, the device must be run with built-in DC power supply.

2.5 The design of this device has mature consideration of security, but operator should never neglect attention to device state and patient's observation. Cut off the power or take off the electrode when necessary to ensure patient's security.

2.6 Please turn off the device and pull out power supply plug before replacing the fuse or cleaning and disinfection. Don't rub the screen with edge tools or sharp materials.

2.7 Keep the device from water, don't use or store it in the place with the air pressure, humidity or temperature over the standard, bad ventilation, or too much dust.

2.8 Do not use the device in the place with flammable anesthetic gases or other flammable chemicals, otherwise there is a danger of explosion or fire.

2.9 Do not use the device in medical hyperbaric oxygen chamber, otherwise there is a danger of explosion or fire.

2.10 This device is not intended to act directly on the human heart. If this device is used with cardiac defibrillator or other electric stimulating devices at the same time, single-use electrode and ECG lead cables with defibrillation-proof function should be selected. It is better not to use this device with other electric stimulating devices at the same time. If it is necessary, there must be professional technician guiding on the scene, and the selected accessories should be designated by our company.

Warning: Do not operate the instrument on parts of human body with wounds, and do not perform measurements on parts with wounds on the surface.

2.11 When the electrocardiograph is used together with a high-frequency electrosurgical knife, the ECG electrode should be kept away from the contact of the electrosurgical knife to prevent burns and burning of the electrode wires caused by high-frequency sparks.

2.12 When the electrocardiograph is used together with a defibrillator, the operator should avoid contact with the patient or the sickbed. The defibrillation electrode should not directly touch the ECG electrode to prevent sparks from burning the device and the patient.

2.13 Please do not use the electrocardiograph in the environment that is interfered by high-power device such as high-voltage cables, X-rays, ultrasonic machines and electrizer, away from

emission sources such as mobile phones.

2.14 When other devices are connected with this ECG instrument, they must be Type I devices which accord with IEC60601-1. Because the total amount of leakage current may hurt patients, the monitoring of leakage current is carried out and taken charge by connect devices.

2.15 Notes related to EMC

The device complies with the safety standards for medical electrical equipment or system electromagnetic compatibility in IEC60601-1-2. Electromagnetic environments exceeding the IEC60601-1-2 standard may cause harmful interference to the device or prevent the device from performing its intended function or degrade its performance. Therefore, if there is a phenomenon that does not match its function during use, be sure to confirm and eliminate adverse effects before continuing to use it. Corresponding precautions for this situation are given in this manual.

- The device or system should not be used near or stacked with other devices. If it must be used near or stacked with other devices, it should be observed and verified that the device is working normally under the configuration it is using.
- In addition to transducers and cables sold by the manufacturer of the device or system as spare parts for internal components, use of accessories and cables outside of the regulations may result in reduced muscle-building emitted by device or system and interference immunity.
- Effect from radiated electromagnetic waves:

The use of a mobile phone may affect the operation of the device. When installing medical electrical equipment, be sure to remind people around the device to turn off mobile phones and small radios.

Effect from shock and conduction electromagnetic waves:

High frequency noise from other equipment can enter the device through the AC socket. Please identify the source of noise, if possible, stop using the equipment. If the equipment can not be deactivated, use noise cancellation equipment or take other measures to reduce the impact.

Effect from static electricity:

Static electricity in a dry environment(indoor) may affect the operation of the device, especially in winter. Before using the device, humidify the indoor air or discharge the static electricity from the cable and ECG record personnel.

Effect from thunder and lightning:

If there is thunder and lightning nearby, it may cause a voltage surge in the device. If you are concerned about danger, pull the AC power plug and use the internal power supply.

2.16 Notes concerning ECG waveform measurement and analysis

2.16.1 P wave and Q wave identify are not always reliable with intensive EMG or AC interference. Neither are the ST segment and T wave with baseline drift.

2.16.2 Winding and unclear end position of S wave and T wave may cause error in measurement. 2.16.3 When R wave is uninspected caused by some leads off or QRS wave low voltage, the heart rate measurement may deviate greatly from the correct.

2.16.4 In case of QRS low voltage, ECG axis calculation and border-point identify of QRS wave

are not always reliable.

2.16.5 Occasionally, frequent ventricular premature complexes may be identified as dominant beat.

2.16.6 Merging of versatile arrhythmia may result in unreliable measurement because of the difficulty in distinguishing P wave in such situation.

2.16.7 The device has an automatic analysis function that automatically analyzes the obtained ECG waveform without reflecting all the patient's status. The results of the analysis may sometimes not comply with the doctor's diagnosis. Therefore, the final conclusion needs to be comprehensively analyzed by doctors in combination with analysis results, patient clinical characterization and other test results.

Chapter3 Warranty Regulation

3.1 In normal use, under strict observance of user manual and operation notes, in case of failure, please contact with our customer service department. Our company has the sales record and customer archives for each device. The customer has one year's warranty service from the date of shipping according to the following conditions. To supply all-around and quick maintenance service for you, please mail the maintenance card to us in time.

3.2 Our company may adopt such ways as guidance, express to company or calling in, etc to carry out warranty promise.

3.3 Even in warranty period, the following repairs are charged in principle.

3.3.1 Faults or injuries caused by misuse not according to user manual and operation notes.

3.3.2 Faults or injuries caused by dropping accidentally when moving after purchasing.

3.3.3 Faults or injuries caused by repair, reconstruction, decomposition, etc not in our company.

3.3.4 Faults or injuries caused by improper storage or force majeure after purchase.

3.3.5 Faults or injuries caused by improper thermal recording paper.

3.4 The warranty period for accessories and fray parts is half a year. Power cable, recording paper, operation manual and packing material are excluded.

3.5 Our company is not responsible for the faults of other connected devices caused by the faults of this device directly or indirectly.

3.6 The warranty will be canceled if we find the protection label has been destroyed.

3.7 For charged maintenance beyond warranty period, our company advises to continue using

"Maintenance contract regulation". Please refer to our customer service department for details.

Chapter4 Working Principle and Structural Characteristics

4.1 Brief Description and Block Diagram of the Working Principle

4.1.1 The power supply unit

(1) Principle of power supply

The switching power supply provides +24V working voltage for the thermal print head, provides constant voltage current limiting charging for the rechargeable lithium battery in the device through the DC-DC circuit, and generates +5V and +12V voltage through the power conversion to supply power to the corresponding modules. At the same time, the lithium battery in the device can independently complete working requirements of each module in the device through the buck-boost circuit.

(2) Principle block diagram is shown in Figure 4-1.

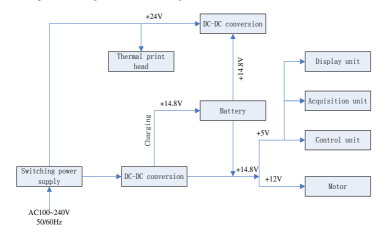


Figure4-1 Block diagram of power principle

Note: The principle block diagram and component list are only available to service stations or maintenance personnel designated by our company.

4.1.2 Signal acquisition unit

The signal acquisition unit uses a floating setting, which is a signal acquisition and processing system, including analog circuit part and A/D conversion and data processing part with sampling accuracy of 24 bits. The analog circuit consists of signal following, amplification, anti-aliasing low-pass filtering, lead-off detection and overload detection. CPU system is responsible for coordinating the work of each circuit such as the A/D converter, the lead-off detection circuit and the overload detection circuit, completes signal acquisition, processing, and lead-off detection. Control information and A/D conversion and data acquisition between the floating circuit and the solid circuit are transmitted through the optoelectronic coupler.

4.1.3 Control unit

(1) Principle of control unit

The control system consists of printing system, button system, liquid crystal display system, and signal acquisition system. The ECG signal sent from the signal acquisition system through the high-speed optoelectronic coupler is received by the CPU system, after digital filtering, gain adjustment and motor drive, it is sent to the printing system to print the ECG waveform. After the printing is completed, the CPU system processes waveform measurement and analysis. The CPU system also receives an interrupt signal from the button system to complete the interrupt processing. In addition, the lead-off signal, paper out detection, battery voltage management, and automatic power-off are also managed by the CPU system. The liquid crystal controller receives data and commands from the CPU system to complete the display of the control state of the device.

(2) Principle block diagram is shown in Figure 4-2.

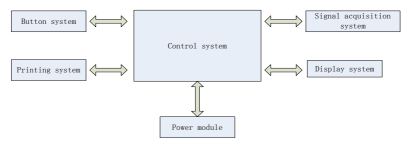


Figure 4-2 Block diagram of control unit

4.2 Name of each part and its function

4.2.1 Front view

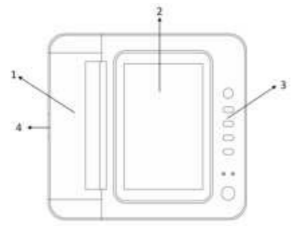


Figure 4-3 Front view

1. Paper compartment cover

Keep the paper compartment closed, hold the printing paper

- 2. Display screen Display patient's ECG and related information
- 3. Button area Control the operations of the device
- 4. Toggle switch

Press down the toggle switch to open the paper compartment cover



- Do not put heavy objects on the screen or hit against it, otherwise the screen will be damaged.
- > If the device is not in use, cover it to prevent liquid spills on the screen.
- Do not use sharp stuff to operate the buttons, otherwise it may case permanent damage to the buttons.
- 4.2.2 Side view

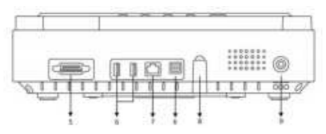


Figure 4-4-1 Side view 1

5. Lead cable interface

Connect with lead cables.

6. USB interface

Communicate with the computer. The ECG data and analysis results can be transmitted to a computer, by using the computer, many functions can be achieved, such as archiving, managing, and analyzing ECG data, facilitating clinical research, organizing teaching and training as well as program upgrade, import/export of cases and connect to network, external printers, etc.

7. Network interface

Connect with the LAN, then perform case analysis and remote control by expert in the LAN

8. Upgrade interface

A USB interface used for program upgrading

9. Equipotential terminal

Connect with the potential equalization conductor

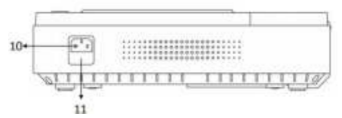


Figure 4-4-2 Side view 2

10.Input socket

Connect with the AC power cable.

11.Fuse

Built-in fuse tube, T3.15AH250V. It can avoid the damage to human body caused by large voltage and large current generated by grid pollution.

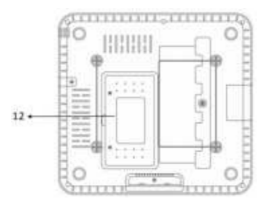


Figure 4-5 Bottom view

12.Battery compartment

Built-in rechargeable lithium battery

4.2.3 Buttons

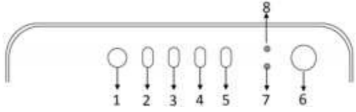


Figure 4-6 Schematic diagram of buttons

1. START/STOP

Used to start/stop the sampling.

2. MODE

When the device in sampling interface, use MODE button to select the print mode.

3. SEN

Used to adjust the sensitivity manually.

4. SPEED

Used to set the ECG recording speed.

5. PRINT

Used to print the sampled ECG waveform or end the printing.

6. ON/OFF

When the device is turned on, short press this button, it will prompt whether to shut down the device, long press this button to turn off the device.

7. Power status indicator

Green indicates that the AC power supply is provided. At this time, there is no battery in the device or the battery is full. Red and green two colors indicate that the battery is being charged.

8. Startup indicator

The indicator lights in green after the device is turned on.

4.2.4 Meaning of symbols

AC 100-240V	Alternating current
\Rightarrow	Equipotential point, the equipotential point of this device is combined with the protective ground.
\wedge	Caution! Refer to the accompanying document.
8	Type CF equipment, with defibrillation-proof function
4	USB interface
매	Network interface
PATIENT	Lead cable socket
T3.15AH250V	Fuse specification
SN	Serial number
***	Manufacturer
\sim	Date of manufacturer
LOT	Batch code

(CATER)	Latex free
Ś	Atmospheric pressure limitation
X	Temperature limitation
ø	Humidity limitation
	Refer to instruction manual/booklet
\bigtriangleup	Indoor use only
Ξ <u>Ι</u>	This way up
[¶]	Fragile, handle with care
Ť	Keep away from rain
	Stacking limit by number
REF	Catalogue number
EC REP	Authorized representative in the European Community
R	Waste disposal symbol. This symbol indicates that electrical and electronic equipment waste cannot be disposed of as unsorted municipal waste and must be recycled separately.
CE	This item is compliant with Directive 93/42/EEC of 14 june 1993 concerning medical devices;Including,at 21 march 2010,the amendments by Council Directive 2007/47/EEC
	General warning sign NOTE : Background col our: yellow Triangular band: Black

Chapter 5 Operation Precautions

5.1 Precautions before use

5.1.1 For safe and effective use, please read the user manual carefully before operation.

5.1.2 Check to ensure that the device is in good condition.

5.1.3 The device shall be placed on a flat surface, and moves gently to avoid strong vibration or shock.

5.1.4 Check to ensure that the lead cables are correctly connected, and the device grounding is correct.

5.1.5 The AC frequency and voltage should comply with the requirements, and enough current capacity should be guaranteed.

5.1.6 When using the battery for power supply, check to ensure that the battery voltage and battery status is in good condition, and the battery has enough power.

5.1.7 When the device is used together with other equipment, all devices and equipment should be equipotential grounded in order to protect the user and operator.

5.1.8 Install the device where easily grounded in the room. Do not allow the patient and patient-connected lead cables and electrodes to come into contact with other conductor parts, including the earth or a hospital bed.

5.1.9 Clean the lead cable with neutral solvent. Do not use alcohol-based cleaners or gemicides.

5.1.10 Ensure that the device is running within the normal ambient temperature range of 5° C to 40°C. If the device is stored at a higher or lower temperature, leave it in the operating environment for approximately 10 minutes before use in order to ensure the normal work.

5.2 Precautions during operating

5.2.1 The printing can be started after the ECG waveform is stable.

5.2.2 During using, the doctor should observe the patient carefully and cannot leave the operating site. If necessary, turn off the power or remove the electrode to ensure patient safety.

5.2.3 The patient and the device can only be connected via lead cables through the electrodes, in order to avoid patient touches other parts of device or conductors.

5.2.4 Patient can not move during operating.

5.2.5 Maintenance or repair to the device or accessory is not allowed during using.

5.3 Precautions after use

5.3.1 Set the states of all functions to initial states.

5.3.2 Cut off the power, gently remove the electrodes and limb clips, then remove the lead cables, do not pull with force.

5.3.3 Clean the device and all accessories, and store them for the next use.

Chapter 6 Preparations before Operation

6.1 Recording paper

6.1.1 The following thermal recording paper can be applied to the device:

Roll paper: 210 mm(W)×20 m(L), 210 mm(W)×30 m(L), 216 mm(W)×20 m(L) (optional); Folding paper: 210×140-20M

ANote: The recording paper should be aligned with the slot of the paper compartment

cover. It is recommended to leave 2cm paper outside.

6.1.2 If the recording paper runs out during recording, the device will stop printing automatically, and the screen will display a prompt of lack of paper.

6.2 Power supply connection

6.2.1 AC

Insert one end of the provided three-core power cord into the device's input socket, and insert the other end into a three-core power socket that meets the requirements. Ensure that the connection is secure and reliable, and the device is automatically grounded.

When the machine is used in conjunction with other medical equipment, use the supplied potential equalization wire to connect the equipotential terminal of the device to the equipotential terminal of the connected equipment to prevent leakage current and protect the device.

6.2.2 Battery

The device has a built-in rechargeable lithium battery, which does not need to be re-installed by the user. Check the battery's power and status before use.

Note: Connect one end of the potential equalization wire to the equipotential terminal of the device, and connect the other end to the ground to enhance the reliability of the grounding. Do not use other pipes as ground wire, otherwise, the patient may be in danger of electric shock.

6.3 Lead cable connection

Connect the lead cable to the lead cable interface on the device, and fasten it to the device with the fixing knobs at both sides of the lead cable in order to prevent bad connection and affecting the detection.

Mote: The lead cable interface can not be used for other purpose except as the input interface of ECG signals.

6.4 Electrode installation

Proper installation of the electrodes is an important part of accurately recording the electrocardiogram. Make sure the electrodes are in good contact. Old and new electrodes or reusable electrodes and disposable electrodes cannot be used at the same time. If different types of electrodes are used together, it will seriously affect the ECG recording. The electrode or lead plug must not touch other object surfaces or conductors, such as metal beds. Please replace them all when updating the electrodes.

6.4.1 Chest electrodes

As shown in Figure 6-1:

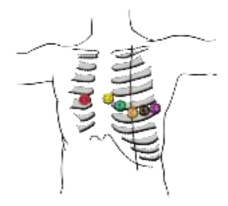


Figure 6-1 Chest electrode installation of E12

The chest electrodes should be installed to the following parts:

C1 (V1) : the fourth intercostal space at the right sternal margin

C2 (V2) : the fourth intercostal space at the left sternal margin

C3 (V3) : between C2 and C4

C4 (V4) : the intersection between midclavicular line and the fifth intercostal space

C5 (V5) : left anterior axillary line on the same plane as C4

C6 (V6) : left midaxillary line on the same plane as C4

Except C1(Vl)~C6(V6), E18M also has the following leads:

Leads attached to right chest C3R(V3R), C4R(V4R), C5R(V5R): symmetric with corresponding left chest leads about the sternum axis

Leads attached to left posterior chest C7(V7), C8(V8), C9(V9): the attachment positions are posterior axillary line of the fifth intercostal space, scapular line, and paravertebral line in turn.

Clean the chest skin where the electrodes to be installed with alcohol, and apply some conductive pastes to these skin (about 25 mm-diameter range) and the edge of the chest electrode suction cup. Squeeze the suction cup to install the chest electrode at proper positions.

Note: The conductive paste coating should be separated from each other, and the chest
electrodes should not touch each other to avoid short circuit.

ANote: Please use qualified conductive paste to avoid damaging the skin.

6.4.2 Limb electrodes

The limb electrodes should be placed on the soft skin of both hands and feet. Before connecting, clean the skin of the electrode installation area with alcohol, and then apply a small amount of conductive paste on the cleaned skin. The electrode connection of the limbs is shown

in Figure 6-2.



Figure 6-2 Installation of limb electrode

6.4.3 Colors of lead cables

As shown in Table 6-1:

	Europe	an standard	American	standard
Electrode position	Mark	Color	Mark	Color
Right arm	R	Red	RA	White
Left arm	L	Yellow	LA	Black
Left leg	F	Green	LL	Red
Right leg	N/RF	Black	RL	Green
Chest 1	Cl	Red	Vl	Red
Chest 2	C2	Yellow	V2	Yellow
Chest 3	C3	Green	V3	Green
Chest 4	C4	Brown	V4	Blue
Chest 5	C5	Black	V5	Orange
Chest 6	C6	Purple	V6	Purple
	С	nly suitable for E18N	1	
Chest 7	C3R	White	V3R	White
Chest 8	C4R	White	V4R	White
Chest 9	C5R	White	V5R	White
Chest 10	C7	White	V7	White
Chest 11	C8	White	V8	White
Chest 12	C9	White	V9	White

Table 6-1 Colors of lead cables

ANote

> It is recommended to install the lead cables after turning off the device.

- > Apply appropriate amount of conductive paste on the electrode when installing the electrode.
- > If the ECG waveform does not appear for a long time, check if the electrode is in

good contact with the skin.

6.4.4 Lead method and system

As shown in Figure 6-3:

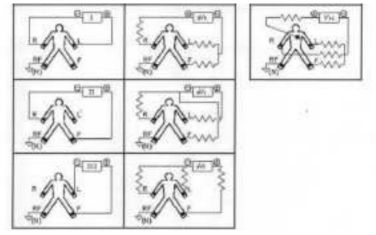


Figure 6-3 E12 Lead system

6.4.5 Lead-off and overload indication

The device can check the connection status of the lead at any time. If lead-off or overload is detected, the screen will display corresponding prompts.

ANote

- > A red lead icon displayed in the status bar under the sampling interface represents lead-off. A yellow lead icon represents overload.
- > When the connection between lead cable and patient/the device is not reliable, and the ECG signal can not correctly transmitted, the device displays lead-off.
- In the printed report, lead-off is marked with "*", and lead overload is marked with "+".

Chapter 7 Operation Instructions and Parameter Setting

7.1 Main Interface

The main interface shows the following information:

♦ Status bar

- Network: the current status of network, including wired network, Wi-Fi
- Battery: the current battery status (refer to 9.1)
- Time: the system time

Functional panel

- Gather: to input the case information, then enter the sampling interface to realize waveform sampling, display and report printing.
- Archive: to enter the case management interface, in this interface, user can query, modify, delete and export case information or review the case to view and print the diagnosis report.
- Last: to quickly modify and review the latest collected case and view its diagnosis report.
- System Setup: to making setting to the system, sampling, print, network, service and time, etc.
- Print Setup: to set the print mode, print style and print content, etc.
- Placement: to view the lead placement schematic diagram.
- About: to view the software version, software establish time, wired network address, wireless network address and used space.

Click the functional module on the screen to quickly set the corresponding function.

7.2 Sampling

Click "Gather" on the main interface or press the START/STOP button to enter the case information input interface.

7.2.1 Input case information

In the case information input interface, input the patient's information by typing or selecting, or obtain patient information via the ID card reader, or click "Get" to extract patient information from stored cases to avoid repeat operation.

◆ Case data

- Name: 0~18 chars
- Sex: Male, Female
- Section: 0~16 chars
- Age: 0~150
- Operator: 0~16 chars
- Bed ID: 0~16 chars
- Room ID: 0~16 chars
- Accession Number: 0~16 chars
- Custom 1: 0~24 chars
- Content of custom 1: 0~24 chars
- Custom 2: 0~24 chars
- Content of custom 2: 0~24 chars

- Custom 3: 0~24 chars
- Content of custom 3: 0~24 chars
- Source: select from clinic, hospital, emergency, checkup, community
- Pace: Whether the patient has a pacemaker.

Operation field

- Get: obtain the case list in case management. Search a patient's information in the list, select the case item, and information of this patient will be automatically added to the edit box in case information input interface. Custom content can be set according to your needs.
- Gather: refer to 7.2.2

In the case information input interface, click any edit box to pop up the keyboard. Click the "Chinese " key to switch between Chinese and English, click the "Chinese " key to switch between numeric keys, lowercase letters and uppercase letters. The "Chinese" is the space key, click it to enter a space; the "Chinese" is the backspace key, click it to delete the last input character. Click the "ENT" key to confirm the entry and exit the interface.

According to the input limitation, after clicking the "ENT", the maximum allowable characters will be displayed in the edit box.

After entering the patient information, click "Gather" to enter the case sampling interface.

7.2.2 Case sampling

The sampling interface of E12 provides several lead display mode, including 3-lead, 6-lead and 12-lead.

The sampling interface of E18M is similar with the interface of E12, the display mode includes 3-lead, 6-lead, 12-lead and 18-lead.

The sampling interface includes Status bar, Display field and Operation field.

♦ Status bar

- HR: current sampled heart rate value
- Lead-off and overload: In demo mode, it displays "Demo Mode". In sampling mode, it displays the detected lead status. A red lead icon represents lead-off. A yellow lead icon represents overload.

Display content	Explanation
Process	It is printing.
Waiting	It is finishing printing.
No Paper.	Lack of paper, user should restart printing after loading paper.
Print Timeout	Communication failure between this system and printing sub-system.
ECG Timeout	Communication failure between this system and sampling sub-system.
Low Power!	Low power, it cannot start printing.
No USB device	No external printer connected, user should restart printing after connected to the external printer.
Gather Time Less	The sampling time is not enough, the print shall be started after

• System status indication:

reaching the required time period.

Display field

- E12: The screen displays sampled 12-lead ECG waveform, by long pressing the waveform, you can switch between 3-lead, 6-lead and 12-lead. You can slide up and down to view each lead.
- E18M: The screen displays sampled 18-lead ECG waveform, by long pressing the waveform, you can switch between 3-lead, 6-lead, 12-lead and 18-lead. You can slide up and down to view each lead.

Operation field

Control the print display mode of the device through the corresponding operation settings.

- Patient: If patient information is not entered before sampling, click this key to pop up the case information input dialogue box to input information.
- Lead: You can choose to display one, some or all of the leads in the waveform display area in the pop-up dialog box. You can also click "Baseline reset" in the dialog box to realize baseline reset.

ANote: When using the baseline reset function, the low frequency part of the waveform may be attenuated.

- Speed: use the SPEED button to switch the speed between 5 mm/s, 6.25 mm/s, 10 mm/s, 12.5 mm/s, 25 mm/s, 50 mm/s and other options.
- Gain: use the SEN button to switch the gain between 2.5 mm/mV, 5 mm/mV, 10 mm/mV, 20 mm/mV,40 mm/mV and other options. The overall gain (sensitivity) can be checked by calibration function.
- Print mode: in print setup, when the data type is set to "After Print", use the MODE button to switch the print mode between Manual, Auto M×N, Auto M×N+1, Auto M×N+2, Auto M×N+3 and Rhythm M. Refer to the print mode in Section 7.5.3 for the value of M and N.
- Print/End print: use the PRINT button to start or end the printing operation.
 - Auto mode: After starting to print, the system automatically prints and stores the real-time ECG waveform. The length is determined by the relevant settings in the print setup. Based on the settings, the automatic analysis data and conclusions are printed, and the system automatically ends printing.
 - Manual mode: After starting to print, user need to switch the lead to print the waveform of different leads, that is, the ECG printed in the manual mode is asynchronous, and the data is not saved. User need to press the PRINT button again when the print needs to be terminated.
 - If lead-off occurs during sampling process, the printed waveform will be marked with "*".
 - * If lead overload occurs during sampling process, the printed waveform will be

marked with "+".

• End sampling: After the device starts sampling, use the START/STOP button to end the sampling, and back to the main interface.

7.3 Case management

In the main interface, click "Archive" to enter the case management interface. This interface shows all the cases stored in the device. User can search the required case by the query function; modify case information and view stored waveform in the "Review"; and delete cases via delete function.

Case management interface includes Case list field and Operation field.

◆ Case list field

- Patient name
- Sampling time
- Age
- Section
- Sex
- Diagnosis result

Operation field

- Query: refer to 7.3.1
- Export: connect the device with a USB flash disk, and export the case into the **Archive** folder in the USB flash disk.
- Delete: delete the selected case (be careful, unrecoverable) or all cases
- Review: refer to 7.3.3
- Close: exit the case management interface

7.3.1 Query

In the case management interface, click "Query" to pop up the search case dialog box. The dialog box includes Case information field and Operation field.

♦ Case information field

- Accession Number: input the accession number of patient
- Name: input patient's name
- Age: input patient age
- Diagnose: input the diagnosis information of the case to be searched

Operation field

- Search: input the query conditions in the search case interface, click "Search", all cases that meet the query conditions will be displayed.
- Close: exit the search interface.

Suggestion: When there are many cases, it would be better to input accurate query conditions to quickly find the case.

7.3.2 Export

In order that the case is not used or known by unauthorized individuals or entities, click "Export" button in the case management interface to open the password input dialog box (initial password: 888888, which can be set in the system setup, see 7.5.1). After inputting the password,

click "OK" to pop up the case export dialog box.

The case export dialog includes Information field and Operation field.

- Information field
 - Choose: to export all cases or export current case
 - File type:
 - Case report: PDF report and image report, PNG, JPEG and BMP are the format of image report.
 - ✤ aECG: case data that conform to the HL7 standard
 - ✤ DAT: case data, self-defined format
 - Orientation: This item is only valid for case report, which determines the generated report is displayed in horizontal or vertical.
 - Directory: the storage path of exported case report or case data
 - Progress bar: indicates the progress rate of exporting

♦ Operation field

- OK: perform the export operation
- Close: exit case export interface

7.3.3 Review

In the case management interface, select a case to be reviewed, click "Review" to enter the case information dialog, which displays the case information, and user can modify the patient information, switch the lead that placed wrong during sampling, and enter the waveform interface to review the sampling process.

The case information dialog includes Case information field and Operation field.

♦ Case information field

• The items are the same wit the items in 7.2.1.

♦ Operation field

- SwitchChannel: If a lead is placed incorrectly during sampling process, click this key to make a correction.
- Review: review the waveform of the selected case, the review interface is similar with the sampling interface.
- Save: user can modify the patient information of the selected case, then click "Save" button to save the modification.
- Close: exit the interface.

Make sure the input information is correct, click "Review" to enter the review interface, which is similar with the sampling interface.

7.4 Latest case

In the main interface, click "Last" to open the latest sampled case, the interface is similar with the sampling interface, user can view waveform of this case and print its report conveniently. Case review interface of E12 is shown as below.

The case review interface includes Display field and Operation field.

♦ Display field

- Filter mode: the filter mode adopted by this case is displayed in the top right corner of the waveform display area.
- If lead-off occurs during sampling process, the reviewed waveform will be marked with "*".
- If lead overload occurs during sampling process, the reviewed waveform will be marked with "+".

Operation field

- Report: displays data information and diagnosis result of the case.
- In this interface, user can use the MODE button to change the print mode.
- In this interface, user can use the PRINT button to print.
 - If lead-off occurs during sampling process, the printed waveform will be marked with "*".
 - If lead overload occurs during sampling process, the printed waveform will be marked with "+".

7.5 System setup

The device related functions can be set in the system setup, which includes the following setting items.

- System setup
- Sample setup
- Print setup
- Network setup
- Server setup
- Time setup

7.5.1 System setup

The optional content of each setting item and its description are shown in the following table:

Item	Options	Explanation	Default
	[OFF]/[1 min]/	If there is no operation after	
	[2 min]/[5 min]/	reaching the set time, screen	
Back-light	[10 min]/[20 min]/	backlight will turn off. If it is	[OFF]
	[30 min]/[60 min],	set to "OFF", the backlight	
	etc	will always keep on.	
		After setting light degree, the	
Light-degree	0%~100%	screen will display different	50%
		backlight strength.	
	[OFF]/[1 min]/	If there is no operation after	
Auto off	[2 min]/[5 min]/	reaching the set time, the	[OFF]
	[10 min]/[20 min]/	system will automatically turn	[OFF]
	[30 min]/[60 min],	off. If it is set to "OFF", the	

		etc	system will always keep on.	
Low Pov	ver	[None]/[Only once]/ [Always]	The system prompts for low battery according to the setting. If "None" is selected, the system will not make any prompt when battery is low.	[Always]
Languag	e	[Chinese]/ [English], etc	Set up the system default language.	[English]
Demo M	ode	[On]/[Off]	Set to on, the system will run in Demo mode; set to off, the system will run in sampling mode	[Off]
K-B Sou	nd	[On]/[Off]	Set to on, the button makes sound while pressing, set to off, there will be no sound	[Off]
XX7 1	Windows Style	[style 1]/ [style 2]	Set the windows style of the system	[style 1]
Windo ws	Font Size	[Small]/[Medium]/ [Large]	Set the font size of the system	[Small]
Style	Background Color	[Dark]/[Light]	Set the background color of system interfaces	[Dark]
	Upgrade	[Upgrade application from U disk]	Upgrade the program	
	Reset	[Factory Reset]	Restore to factory setup	
More	Log	[Export log to U disk]	Export the running log of the program to U-disk	
	Print Test		Test the print function	
	Temperatur e		Display current temperature of the CPU	
	Password	Within 6 chars	Input the password	[888888]
Default		[Reset this page]	All above settings will restore to default after clicking this button.	

7.5.2 Sample setup

The optional content of each setting item and its description are shown in the following table:

Item	Options	Explanation	Default
AC Filter Enable	[On]/[Off]	Turn on or turn off the AC Filter	[On]
EMG Filter Enable	[On]/[Off]	Turn on or turn off the EMG Filter	[On]

DFT Filter	Enable	[On]/[Off]	Turn on or turn off the DFT Filter	[On]
Low-pass Enable	Filter	[On]/[Off]	Turn on or turn off the Low-pass Filter	[Off]
AC Filter		[50Hz]/[60Hz]	Set the parameter of the AC Filter	[50Hz]
EMG Filte	er	[25Hz]/[30Hz]]/ [35Hz]]/[40Hz]]/ [45Hz], etc	Set the parameter of the EMG Filter	[25Hz]
DFT Filter	-	[0.05Hz]/[0.50H z]/[1.00Hz]/[0.15 Hz]/[0.25Hz]/[0. 32Hz]/[0.67Hz]/ [0.01Hz], etc	Set the parameter of the DFT Filter	[0.50Hz]
Low-pass	Filter	[75Hz]/[100Hz]]/ [150Hz]], etc	Set the parameter of the Low-pass Filter	[75Hz]
Pace Lead		[I]/[II]/[III]/[aVR]/[aVL]/[aVF]/[V 1]/[V2]/[V3]/[V4]/[V5]/[V6]	Set pacing detection lead	[11]
Background Grid		[On]/[Off]	Set to use the background grid or not	[Off]
Sort Lead		[Routine Lead]/ [Cabrera Lead]	Set the arrangement of the leads.	[Routine Lead]
Save Time		[8sec]/[10sec]/[1 5sec]/[20sec]/[25 sec]/[30sec]/[40s ec]/[50sec]/[60se c]/[90sec]/[120se c]/[180sec]/[240s ec]/[300sec]/[360 sec], etc	Set the time length of save data	[10sec]
Save Data Type		[BeginData]/ [After Data]	Set to save the data before clicking the PRINT button, or after clicking	[After Data]
	Heartbeat Sound	[On/][Off]	Turn on or tun off the heartbeat sound	[Off]
Analysi s Set	Premature	1-99	The system will use the input value as a standard of judging premature beat.	78
	Pause	1200-3000	The system will use the input	2000

	Time (ms)		value as a standard of judging beat pause.	
	Tachycard ia (bpm)	80-250	The system will use the input value as a standard of judging tachycardia.	100
	Bradycard ia (bpm)	30-80	The system will use the input value as a standard of judging bradycardia.	60
Default		[Reset this page]	All above settings will restore to default after clicking this button.	

7.5.3 Print setup

The optional content of each setting item and its description are shown in the following table:

Ite	em	Options	Explanation	Default
Print Mode		[Manual]/[Auto M×N]/[AutoM× N+1]/[AutoM×N +2]/[AutoM×N+ 3]/[RhythmM], etc	The system takes the selected option as default print mode. "M" is the strip number, its value range of E12 is 3-12, and E18M is 3-18, refer to the setting of strip number.	[AutoM×N]
Strip	E12	3-12	It represents the number of	[12]
Num	E18M	3-18	channels.	[18]
Speed		[5mm/s]/[6.25m m/s]/[10mm/s]/[1 2.5mm/s]/[25mm /s]/[50mm/s], etc	Set the waveform speed displayed on the screen.	[25mm/s]
Gain		[1.25mm/mV]/[2 .5mm/mV]/[5m m/mV]/[10mm/ mV]/[20mm/mV]/[40mm/mV]/[1 0/5mm/mV]/[20/ 10mm/mV], etc	Set the gain of ECG	[10mm/mV]
Auto Strip		[2.5sec]/[3sec]/[4 sec]/[5sec]/[6sec] /[8sec]/[10sec]/[1 5sec]/[20sec]/[25 sec]/[30sec], etc	The system takes the selected option as the print time length of each strip.	[2.5sec]
Print Width		[1]/[2]/[3]/[4]	Set the print width	[1]
Phase Mode		[Continuity]/[In- phase]	The display mode of multiple rows waveform	[Continuity]
Data Type		[Print Immediately/	Set to print immediately or after reviewing. "Print	[Print Cache]

			[Print Cache]	immediately" does not allow]
				5	
				repeat sampling; "Print cache" allows the device to take more	
				sampling, but it will re-timing.	
				The option selected will be	
				used as printing gain mode.	
				"Smart" means the system will	
Lead C	bain		[Smart]/[Current]	adjust gain automatically to fit	[Smart]
				paper height; "Current" means	
				it will use screen waveform	
			gain as that of printing.		
			[I]/[II]/[III]/[aVR		
		E12]/[aVL]/[aVF]/[V		
		212	1]/[V2]/[V3]/[V4	The main lead when print	
]/[V5]/[V6]	mode is "Auto M×N+1",	
	Rhyt		[I]/[II]/[III]/[aVR	"Auto M×N+2" or "Auto	[II]
	hm1]/[aVL]/[aVF]/[V	$M \times N+3$ "; the main lead when	[11]
		E18	1]/[V2]/[V3]/[V4	print mode is "Rhythm M".	
		М]/[V5]/[V6]/[V3		
			R]/[V4R]/[V5R]/		
			[V7]/[V8]/[V9]		
			[I]/[II]/[III]/[aVR		
	E10	E12]/[aVL]/[aVF]/[V		
		LIZ	1]/[V2]/[V3]/[V4		
]/[V5]/[V6]	The secondary main lead when	
Rhyt	Rhyt		[I]/[II]/[III]/[aVR	print mode is "Auto M×N+2"	FV/11
hm	hm2]/[aVL]/[aVF]/[V	or "Auto $M \times N+3$ ".	[V1]
		E18	1]/[V2]/[V3]/[V4	of Auto MAN+5 .	
		М]/[V5]/[V6]/[V3		
			R]/[V4R]/[V5R]/		
			[V7]/[V8]/[V9]		
			[I]/[II]/[III]/[aVR		
		E12]/[aVL]/[aVF]/[V		
		E12	1]/[V2]/[V3]/[V4		
]/[V5]/[V6]		
	Rhyt		[I]/[II]/[III]/[aVR	The third main lead when print	[3/0]
	hm3]/[aVL]/[aVF]/[V	mode is "Auto M×N+3".	[V2]
		E18	1]/[V2]/[V3]/[V4		
		М]/[V5]/[V6]/[V3		
			R]/[V4R]/[V5R]/		
			[V7]/[V8]/[V9]		
			Set to automatically active the	10.07	
Arrhythmia		[On]/[Off]	print function when	[Off]	
		1	1		

			arrhythmia occurs during ECG	
			sampling process.	
Period		[OFF]/ [per 1 min]/ [per 2 min]/ [per 5 min]/ [per 10 min]/ [per 20 min]/ [per 30 min]/ [per 60 min], etc	During the ECG acquisition process, the system will automatically activate the printing operation according to the selected time interval. When the printing mode is manual mode, the printing will output "Auto 12×1" format, otherwise, it will output according to the current setting mode.	[OFF]
Printer set	Print Device	[Inside]/[Outside A4]	Choose to print out the ECG waveform by thermal printing system or USB external printer	[Inside]
	Paper type	[Roll paper]/ [Folding Paper]	Set the thermal paper type	[Roll paper]
	Orientatio n	[Portrait]/ [Landscape]	Set the print direction of external USB printer	[Portrait]
	Info Align	[Right]/[Top]	Set the position of patient information for thermal print	[Right]
	Print Grid	[On]/[Off]	Set to print the background grid or not	[Off]
	Print time mark	[On]/[Off]	Set to print the time mark or not	[Off]
Print Content	MultiSign al	[On]/[Off]	Set to print the multiple signal or not	[Off]
	Average QRS	[On]/[Off]	Set to print the average QRS or not	[On]
	Diagnosis Conclusio n	[On]/[Off]	Set to print the diagnosis conclusion or not	[On]
	Minnesot a Code	[On]/[Off]	Set whether the Minnesota code is displayed in the printed diagnosis conclusion	[Off]
	Diagnosis Data	[On]/[Off]	Set to print the diagnosis data or not	[On]
	Hospital	0-64 chars	Input the name of hospital printed in the report	Blank
Default		[Reset this page]	All above settings will restore to default after clicking this button.	

ANote: The auto strip, average QRS, print diagnosis and period are only optional in auto mode and rhythm mode.

7.5.4 Network setup

The optional content of each setting item and its description are shown in the following table. When the DHCP is turned to On state, the static IP and other items are unable to use.

Item		Options	Explanation	Default
Wi-Fi		[On]/[Off]	Turn on or turn off the Wi-Fi	[On]
Wired		[On]/[Off]	Turn on or turn off the wired network	[On]
	DHCP	[On]/[Off]	Turn on or turn off the DHCP	[On]
	IP	0-64 chars	Input IP address	Blank
	Subnet	0-64 chars	Input the subnet mask	Blank
Wi-Fi		0-64 chars	Input the default gateway	Blank
		0-64 chars	Input the DNS server	Blank
	Wifi List	Searched Wi-Fi	Select the network you want to connect to in the Wi-Fi list	None
	DHCP	[On]/[Off]	Turn on or turn off the DHCP	[On]
XX7' 1	IP	0-64 chars	Input IP address	Blank
Wired	Subnet	0-64 chars	Input the subnet mask	Blank
Set	Set Gateway 0-64 chars	0-64 chars	Input the default gateway	Blank
	DNS	0-64 chars	Input the DNS server	Blank
Default		[Reset this page]	All above settings will restore to default after clicking this button.	

7.5.5 Server setup

The optional content of each setting item and its description are shown in the following table:

Item	Options	Explanation	Default
SYNC Mode	[USB]	Set the synchronous mode	[USB]
Default	[Reset this page]	All above settings will restore to default after clicking this button.	

ANote: In addition, we provide network data exchange interface. If necessary, please contact the supplier for specific settings and instructions.

Item Options	Explanation
--------------	-------------

Case Get Mode	[Case From Local]/ [Case From HIS]/ [Case From CallSys]/ [Case From Third-Part]	We provide patient information interface to obtain case information.
Platform	[PHMS platform] /[Third-Part platform] /[FTP]/[Nonuse]	If you need to exchange data, you can choose the corresponding platform, and [Nonuse] does not use the platform.

7.5.6 Time setup

The optional content of each setting item and its description are shown in the following table:

	Item	Options	Explanation	Default	
Adjustment Time		This option is unavailable if "Net Time" is selected.	Adjust the current date and time manually.	Current date and time	
Net Tim	e	[On]/[Off]	Turn on or turn off the network time	[On]	
Time	Server	0-64 chars	Input the server address corresponding to the time zone	cn.ntp.org.c n	
Server	Port	0-64 chars	4 chars Set the server port 123		
	Time Zone	Time zone list	Select the corresponding time zone from the list	[East Eight]	
Default		[Reset this page]	All above settings will restore to default after clicking this button.		

7.6 Print setup

In the main interface, click "Print Setup" to enter the print setup directly.

7.7 Lead placement

In the main interface, click "Placement" to view the schematic diagram of lead placement, or you can refer to Section 6.4 for the connection of electrodes.

7.8 About

In the main interface, click "About" to view the information about the device, which including the following content:

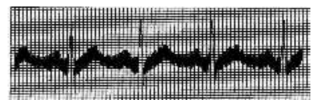
- AppVersion: the version number of current software
- AppBuild: the creation time of current software
- Wired Mac: the MAC address of wired LAN
- Wi-Fi Mac: the MAC address of wireless LAN
- Spaced Used: the percentage of the memory that used in the system

Chapter 8 Troubleshooting

8.1 Auto shutdown

- > The battery is almost running out, which causes overdischarge protection circuit action.
- The voltage of AC power supply is too high, which causes overvoltage protection circuit action.

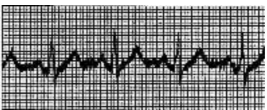
8.2 AC interference



- > Whether the device is grounded reliably?
- > Whether the electrode or lead cable is connected correctly?
- Whether the electrodes and skin are daubed with enough conductive paste?.
- > Whether the metal bed is grounded reliably?
- > Whether the patient is touching the wall or metal parts of the bed?
- > Whether the patient touches other people?
- Whether there is high power electric equipment working nearby? Such as X-ray machine or ultrasonic device, etc.

Note: If the interference can not be removed after taking above measures, please use a AC filter.

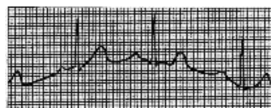
8.3 EMG interference



- ➤ Whether the room is comfortable?
- ➤ Whether the patient is nervous?
- ➤ Whether the bed space is narrow?
- ➤ Whether patient speaks during recording?
- ➤ Whether the limb electrode is too tight?

Mote: If the interference can not be removed after taking above measures, please use a EMG filter. The ECG waveform recorded at this time will be slightly attenuated.

8.4 Baseline drift



- > Whether the electrode installation is stable?
- Whether the connection of lead cables or electrodes is reliable?
- ➢ Whether the electrodes and patient skin are cleaned and are daubed with enough conductive paste?
- > Whether it is caused by patient's movement or breathing?
- > Whether the electrodes or leads are in bad connection?

ANote: If the interference can not be removed after taking above measures, please use a baseline filter.

8.5 Troubleshooting list

Phenomenon	Cause of failure	Solutions
Too large interference, disorderly waveform	 Grounding cable is not connected reliably. Lead cables are not connected reliably. There is AC interference. Patient is nervous and can not keep quiet. 	 Check the power cord and lead cables. Let the patient prepare for the measurement.
Baseline burr	 AC interference is large. Patient nervous, and EMG interference is large. 	 Improve the environment. If the bed is made of steel, replace it. The power cable and lead cables are not parallel or too close to each other.
Not regular waveform, large up-and-down, beeline figure	 Bad electrode conductivity. Low battery. Bad connection between electrodes and patient skin. Loose connection between lead cables and the device's plug. Bad connection between electrodes and lead cables. 	 Use alcohol of high quality. Clean electrode slice and the skin under the electrode with alcohol. Charge the battery.

Baseline draft	 Low power. Patient movement. 	 Charge the battery. Keep patient still.
Unclear waveform	 Low battery. The printer head surface is dirty. The thermal paper problem. 	 Charge the battery. Cut off the power, clean the printer head with alcohol, air dry. Replace the thermal print paper with specified one.

Chapter 9 Maintenance

9.1 Battery

9.1.1 The device is designed with built-in full-sealed and maintenance-free rechargeable lithium battery, also equipped with perfect auto-charging-discharging monitor system. When the device is connected to AC power supply, the battery will be charged automatically. Battery status will be displayed on right edge of LCD screen in powering on state. After absolutely discharged, the battery needs 3.5 hours to charge to 90%, and 5 hours to charge to full capacity.

No.	Icon	Description
а	÷	Using AC power supply, and the battery is full or no battery in the device
b	ļ	Using battery, and battery is full
с		Using battery, and battery level is 3/4 of battery full
d	Ę	Using battery, and battery level is 1/2 of battery full
е		Using battery, and battery level is 1/4 of battery full
f		Using battery, and the battery is low. It is recommended to charge the battery and use AC power supply; or the battery status is unknown, this happens at the starting of turning on the device.
g	1	Charging the battery
h		Battery is charged to full
i	. 13351	Battery overheat

ANote: When charging the battery, the displayed status of battery level switches between

icon b to icon e.

9.1.2 The device can continuously print for 3 hours or work for more than 10 hours in standby mode when battery is completely charged. When the device is powered by battery, a battery icon will be displayed on the LCD screen, showing the battery capacity in 5 modes. When the battery capacity is too low for the device to operate, the device will turn off automatically to avoid permanent damage to the battery.

Note: The above data is obtained by printing demo waveform under the test environment of temperature 25 °C, speed 25mm/s and gain 10mm/mV. In actual use, the operation time may be shorten due to operation condition and environment.

9.1.3 The battery should be recharged in time after discharged completely. If not used for long period, the battery should be recharged every 3 months, which can extend the life of the battery.

9.1.4 When the battery can not be recharged or works no more than 10 minutes after fully charged, please replace the battery.

ANote

- Do not try to dismantle the sealed battery without permission. The replacement of battery shall be carried out by professional maintenance personal authorized by our company, and the same model of rechargeable battery provided by our company should be used.
- Do not touch the positive and negative terminals of the battery directly with wire, otherwise there is a danger of fire.
- Do not use the battery near fire sources or in environments where the temperature exceeds 60°C. Do not heat the battery or throw it into fire, water and avoid splashed by water.
- Do not puncture, hammer or strike the battery or destroy it by other ways, otherwise it will cause battery overheat, smoke, deform or burn dangers.
- Keep away from the battery when it appears leakage or emitting unpleasant smell. If the battery electrolyte leaks onto the skin or clothes, clean with water immediately. If the electrolyte accidentally enters your eyes, do not rub your eyes, immediately clean with water and see a doctor.
- If the battery reaches its service life, or battery smell, deform, discolor or distorted appears, please stop using the battery and dispose it in accordance with local regulations.

9.2 Recording paper

In order to ensure the quality of the ECG waveform, please use the high-speed thermal recording paper supplied or specified by the company. If you use unspecified recording paper, the recorded ECG waveform may be blurred, faded, and the paper feeding may not be smooth. This may even increase the wear of the device and shorten the service life of important parts such as the thermal print head. For information on how to purchase such recording paper, please contact your dealer or the company. Please be careful!

9.2.1 When using recording paper, it is absolutely not allowed to use recording paper with wax on the surface or in grayish/black color. Otherwise, the wax will stick to the heating part of the print head, resulting in abnormal work or damage of the print head.

9.2.2 High temperature, humidity and sunlight may cause the recording paper to change color. Please keep the recording paper in a dry and cool place.

9.2.3 Please do not place the recording paper under fluorescent light for a long time, otherwise it will affect the recording effect.

9.2.4 Please do not to put the recording paper together with the PVC plastic, otherwise the color of recording paper will change.

9.2.5 Please use the recording paper with specified dimension. Recording paper that does not meet the requirements may damage the thermal print head or silicone rubber roller.

9.3 Maintenance after use

9.3.1 Press the power button to shutdown the device.

9.3.2 Unplug the power cord and lead cables. Hold the header of plug to disconnect, and do not

pull the cable with force directly.

9.3.3Clean the device and accessories, cover them up to against dust.

9.3.4 Store the device in a cool and dry place, avoid strong vibration when moving.

9.3.5 When cleaning the device, do not immerse it in the cleaner. Power supply must be cut off before cleaning. Use neutral detergents for cleaning. Do not use any detergent or disinfectant containing alcohol.

9.4 Lead cables and electrodes

9.4.1 The connectivity of the lead cable can be detected by the multimeter. Check whether each wire of the lead cable is in good contact according to the following table. The resistance of each wire from the electrode plug to the corresponding pin in the lead cable plug should be less than 10Ω . The integrity of the lead cable must be checked regularly. Any lead wire damage will cause a false waveform of the corresponding lead or all leads on the ECG. The lead cable cable can be cleaned with neutral solvent. Do not use the detergent or germicide containing alcohol (Please do not immerse the lead cables in liquid for cleaning).

^ΔNote: The resistance of lead cable with defibrillation-proof function protection function is about 10KΩ.

Mark	L	R	C1	C2	C3	C4	C5	C6	F	N
Pin position	10	9	12	1	2	3	4	5	11	14

Table 10 Lead cable mark and pin position table

9.4.2 Bending or knotting will shorten the service life of the lead cable. When using it, please straighten the lead cable first.

9.4.3 The electrode should be well stored. After long time use, the surface of the electrode may oxidize and discolor due to corrosion and other factors, which may affect the signal acquisition. In this case, the electrode must be replaced.

9.5 Silicone rubber roller

The silicone rubber roller should be smooth and free of stains, otherwise it will affect the ECG recording effect. In order to remove the stains on the roller, please use a clean soft cloth damped with a small amount of alcohol to wipe it along the longitudinal direction, and scroll the roller in the paper conveying direction while wiping until it is clean.

9.6 Cleaning of thermal print head

Dirt and dust on the surface of the TPH can affect the clarity of the waveform. To clean the print head surface, open the paper compartment cover after turning off the device, use a clean and soft cloth dampened with alcohol to wipe the surface gently. For the residual stains on print head, moist it with a little alcohol first, then wipe with a soft cloth. Never use hard objects to scratch the surface, otherwise the print head will be damaged. Wait until the alcohol has evaporated, then close the paper compartment cover. The print head should be cleaned at least once a month during normal use.

9.7 Fuse replacement

Unplug the power cord, pull out the fuse box and replace the fuse, the specification of fuse is T3.15AH250V, as shown in Figure 9-1:



Figure 9-1 Replacing the fuse



If the fuse blows again after replacing a fuse of the same specification, the device may exists other problems, please cut off the power supply and contact the after-sales service of our company or designated service center.

9.8 Disposal of product scrap

The disposal of packaging materials, waste battery and end-of-life device should obey the local laws and regulations, and user should treat the scrapped products and materials properly according to the laws and regulations, and try to support the classification and recycling work.

9.9 Others

9.9.1 Do not open the device enclosure to avoid electric shock danger.

9.9.2 The device associated circuit schematics and critical parts list are only available to authorized service station or maintenance personnel, who is responsible for maintenance of the device.

9.9.3 The device belongs to measuring instrument. User should send the device to national designated inspection institution for inspection according to the requirements of the national metrological verification procedure. The device shall be inspected at least once per year, and all the accessories should be inspected and maintained at least once every six months.

Chapter 10 Packing List and Accessories

10.1 Accompanying accessories

When the device is shipped from the factory, the intact packaging should contain the following contents, as shown in Table 11:

Name	Quantity
Electrocardiograph	1 pc
Chest electrodes (suction cup/electrode slice)	E12 1set (6 pcs)
	E18M 2set (12 pcs)
Limb electrodes (limb clip)	1 set (4 pcs)
ECG lead cable	1 pc
Potential equalization wire	1 pc
Power cord	1pc
User manual	1 pc
Recording paper	1 pc

Table	11	Packing	list and	accessories
raoic	11	1 acking	mot and	accessories

10.2 Notes

10.2.1 Please follow the instructions on the package when opening the package.

10.2.2 After unpacking, please check the accessories and accompanying documents in accordance with the packing list, then start inspecting the device.

10.2.3 If the packaging content does not meet the requirement or the device does not work properly, please contact our company immediately.

10.2.4 Please use the accessories provided by our company, otherwise the performance and safety of the device may be affected. If accessories provided by other company need to be used, please first consult the after-sales service of our company, or we will not responsible for any caused damages.

10.2.5 The package shall be kept properly for future use in regular maintenance or device repair.

Appendix I ECG Automated Measurement & Interpretation Guide

1. Preface

The appendix describes the functions of ECG automated measurement and automated interpretation. It explains the specific implementation method, algorithm and formulas related to these two functions, as well as the content output by the automated measurement and automated interpretation.

According to the requirement of IEC60601-2-51:2003 Medical electrical equipment - Part 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiographs, Clause 50 Accuracy of operating data, the appendix gives a description of verification process and results of the performance for automated measurement and automated interpretation.

The appendix also contains rhythm diagnosis function, which interprets the ECG database used for rhythm diagnosis and accuracy verification results of rhythm diagnosis.

2. Automated measurement parameters and Automated interpretation items

The output measurement parameter, interpretation item and others that require explanation are as follows:

No.	Parameter	Unit
1	HR	bpm
2	PR-interval	ms
3	P-duration	ms
4	QRS-duration	ms
5	T-duration	ms
6	QT/QTc	ms
7	P/QRS/T electric axis	deg
8	R(V5)/S(V1)	mV
9	R(V5)+S(V1)	mV

2.1 Measurement parameters

2.2 Interpretation items

No.	Item
1	No abnormal
2	Sinus mode Bradycardia
3	Sinus mode Tachycardia
4	Left atrium Hypertrophy
5	Right atrium Hypertrophy
6	Dual atrium Hypertrophy
7	QRS low voltage
8	Cardiac electric axis normal

9	Left axis deviation	
10	Right axis deviation	
11	Completeness Right Bundle branch block	
12	Completeness Left Bundle branch block	
13	No Completeness Right Bundle branch block	
14	No Completeness Left Bundle branch block	
15	V1 shows RSR' type	
16	Left anterior fascicular block	
17	Left posterior fascicular block	
18	Left ventricular hypertrophy	
19	Right ventricular hypertrophy	
20	I atrioventricular block	
21	Early anteroseptal MI	
22	Possible acute forepart anteroseptal MI	
23	Old anteroseptal MI	
24	Early anterior MI	
25	Possible acute anterior MI	
26	Old anterior MI	
27	Early extensive anterior MI	
28	Possible acute extensive anterior MI	
29	Old extensive anterior MI	
30	Early apical MI	
31	Acute apical MI	
32	Old apical MI	
33	Early anterolateral MI	
34	Possible acute anterolateral MI	
35	Old anterolateral MI	
36	Early high lateral MI	
37	Possible acute high lateral MI	
38	Old high lateral MI	
39	Early inferior MI	
40	Possible acute inferior MI	
41	Old inferior MI	
42	Early inferolateral MI	
43	Possible acute inferolateral MI	
44	Old inferolateral MI	
45	ST depression, mild anteroseptal myocardial ischemia	

46	ST depression, mild anterior myocardial ischemia
47	ST depression, mild extensive anterior myocardial ischemia
48	ST depression, mild apical myocardial ischemia
49	ST depression, mild anterolateral myocardial ischemia
50	ST depression, mild high lateral myocardial ischemia
51	ST depression, mild inferior myocardial ischemia
52	ST depression, mild inferolateral myocardial ischemia
53	ST depression, anteroseptal myocardial ischemia
54	ST depression, anterior myocardial ischemia
55	ST depression, extensive anterior myocardial ischemia
56	ST depression, apical myocardial ischemia
57	ST depression, anterolateral myocardial ischemia
58	ST depression, high lateral myocardial ischemia
59	ST depression, inferior myocardial ischemia
60	ST depression, inferolateral myocardial ischemia

2.3 Intended use

The intended use of the Automated Measurement&Interpretation function is shown as below:

Application and diagnosis	To detect the abnormal of heart of human body, examination items refer to above description
Population	Teenagers and adults, age range: 12-87
Application site	hospitals
Accuracy	The accuracy of this function is reflected by the balance performance of sensitivity and specificity.
Others	This function does not generate any alarm when using, so it should be operated by professional or trained personal.

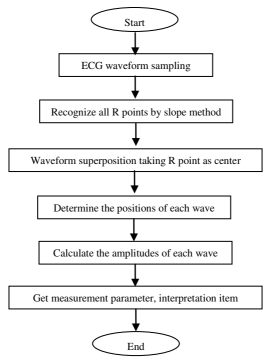
3. Algorithm description

This section describes the algorithm, formulas and judgment conditions for interpretation items related to functions of ECG automated measurement and automated interpretation.

The 12-lead sync ECG waveform passes through the filter (AC, EMG, DFT (if has, and open)) into the module of automated measurement and automated interpretation.

The module of automated measurement and automated interpretation mainly includes process of find the cardiac impulse location, find the beginning/end for each wave, amplitude calculation, parameters calculation, and interpretations judgment based on known parameters.

The workflow is shown as below:



3.1 Find the cardiac impulse location

1) Data preprocessing, obtain the absolute value trend of slope for each lead; then superimpose each absolute value, obtain the superimposed graph of absolute value of slope.

2) Smoothing filter the superimposed graph on average of width 80ms, obtain the analytical data source DDD.

3) Find the cardiac impulse location, give an initial threshold for searching, orderly scan the data in the analytical data source DDD, then compare it with the threshold value:

When the value is greater than the threshold, it may be the beginning of qrs-complex. If the distance from the previous qrs-complex to the current location is less than 150ms, then give up the location.

Otherwise, take the 1/4 of threshold value as a reference, find the beginning of qrs-complex within 100ms before the current location.

When the value is less than the threshold value, it may be the end of qrs-complex. Take the 1/4 of threshold value as a reference, find the end of qrs-complex.

If the found qrs-complex is wide, this qrs-complex shall be excluded. Otherwise, save the found qrs-complex.

4) Locate: after found the qrs-complex, search the max value point between the beginning point and end point in the ecg original data, mark the point as cardiac impulse location.

5) Dynamically threshold adjustment: after found the cardiac impulse location, use the value at the cardiac impulse location for the dynamically adaptive adjustment of the threshold value. Define the threshold value as 1/3 of the average of the nearest three cardiac impulses.

6) After found the cardiac impulse location, compute the RR-interval and accumulate it with the previous RR-intervals, then count the number of accumulated RR-intervals.

7) Continue searching until the end of data, and calculate the global average value for RR-intervals at the same time.

3.2 Find the beginning/end for each wave

The beginning/end of qrs-complex has been approached in above cardiac impulse locating process, but it is mainly in order to assist to find the cardiac impulse location; in addition, the location is searched based on the slope threshold value, which is imprecise. Here, according to the found cardiac impulse location, the beginning/end of qrs-complex will be sought accurately. Name the cardiac impulse location as the peak of R-wave.

1. Read data

1) Read one data of qrs-complex: take the peak of R-wave as reference, locate directly to the original ecg file, read a piece of data containing the qrs-complex.

2) Preprocessing: superimpose the absolute value of slope for 12-lead signals.

3) Use the preprocessed data to carry on the searching of QRS-complex, P-wave and T-wave as the followings.

4) Read the next data of qrs-complex, repeat step 2 and step 3 until the analyzing of all qrs-complex are finished.

2. Find QRS-complex

1) Calculate the threshold value of S-wave: search the minimal value within 200ms after the peak of R-wave, take the value that equals to minimal value plus 0.4, as the threshold value for finding the end of S-wave.

2) Find the beginning of Q-wave: take 0.5 as the threshold vale, search forwardly starting from R-wave, a point that less than the threshold value, within 0ms-200ms before the peak of R-wave, which is the beginning of Q-wave.

3) Find the end of S-wave: search backwardly starting from R-wave, a point that less than the threshold value of the end of S-wave, within 0ms-200ms after the peak of R-wave, which is the end of S-wave.

3. Find P-wave

1) Peak of P-wave: search the max value within 30ms-100ms before the beginning of Q-wave, temporarily mark the point as the peak of P-wave.

2) Find the end of P-wave: search the minimal value between the peak of P-wave and the beginning of Q-wave, the minimal value plus 0.05 is the threshold value, use the threshold value to find the end of P-wave.

3) Find the beginning of P-wave: search the minimal value within 150ms before the peak of P-wave, the minimal value plus 0.06 is the threshold value, use the threshold value to find the beginning of P-wave.

4) If the found P-wave is narrow, research the P-wave according to the following steps.

5) Change the searching range of 30ms-100ms to 100ms-350ms in step 1, repeat step 1-4.

6) If the found P-wave is still narrow, it means that P-wave doesn't exist.

4. Find T-wave

1) Peak of T-wave: search the max value within 30ms-300ms after the end of QRS-complex, save it as the peak of T-wave.

2) Threshold value of the beginning of T-wave: search the minimal value within 0ms-100ms after the end of QRS-complex, the minimal value plus 1/10 of the peak value of T-wave is the threshold for finding the beginning of T-wave.

3) Threshold value of the end of T-wave: search the minimal value within 200ms after the peak of T-wave, the minimal value plus 1/10 of the peak value of T-wave is the threshold for finding the end of T-wave.

4) Find the beginning of T-wave: in the range between the minimal value in step2 and the peak of T-wave, find a point that less than the threshold value of the beginning of T-wave, the point is the beginning of T-wave.

5) Find the end of T-wave: in the range between the minimal value in step3 and the peak of T-wave, find a point that less than the threshold value of the end of T-wave, the point is the end of T-wave.

5. Explanation of equipotential segment

In searching the QRS-complex, this algorithm adopts the analysis method of superposition of the slopes for all leads, therefore, the equipotential segments before and after the QRS-complex are partly included in the start and end points of the QRS-complex. It is depends on the number of leads containing equipotential segments. If there are more leads containing equipotential segments, the slope value will be smaller after superposition, so it is difficult to meet the threshold condition, and only a small part of the equipotential segments is counted to the start and end points of the QRS-complex. On the contrary, if there are less leads containing equipotential segments, a large part of the equipotential segments will be counted to the start and end points of the QRS-complex. Anyway, the equipotential segments before and after the QRS-complex are partly included in the QRS-complex duration.

3.3 Amplitude measurement

After finding the position of each wave, i.e. the start and end points of P wave, QRS complex and T wave, use the following method to measure P, Q, R, S, ST and T waves of each lead.

1. P-wave

Calculate the average value of the data 20ms before the start point of P wave, and use this average value as the baseline of P wave. Find the max value between the start point and end point of P wave, the difference between the max value and the baseline would be the amplitude of P wave.

2. Q/R/S wave

Calculate the average value of the data 10-30ms before the start point of QRS complex, and use this average value as the baseline of QRS complex. Search boundary points that exceeding the baseline from the start point of Q wave to the end point of S wave. Each adjacent two boundary

points forms a sub-wave. Determine whether each sub-wave is a recognizable minimum wave (see the definition below). If it is a recognizable minimum wave, first identify its direction. If it is above the QRS baseline, it is R wave, if it is below the baseline, it is Q wave or S wave. Find the extreme value of this wave, and the difference between the extreme value and the baseline is the amplitude of Q/R/S wave.

ANote: If there is only one downward wave, its amplitude should be respectively recorded in the amplitude of Q wave and S wave.

3. ST segment

Take above baseline of QRS complex as the ST baseline. Calculate the differences between the ST baseline and the points at 40ms and 60ms after the end point of QRS complex, and calculate the average value of these two differences, the average value is the amplitude of ST segment.

4. T-wave

Calculate the average value of the data 20-50ms after the end point of T wave, and average this value with the QRS baseline in 2, then use the result as the baseline of T wave. Find the max value between the start point and end point of T wave, the difference between the max value and the baseline would be the amplitude of T wave.

5. Recognition of minimum wave

The minimum wave can be recognized by the algorithm according to the requirement of IEC60601-2-51:2003 Medical electrical equipment - Part 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiographs, Annex GG, Clause GG.5 Definition of waveforms, measurement of minimum waves. The wave that meet the following conditions is the minimum wave that can be recognized by the algorithm.

1) The signal part under consideration shows clearly two opposite slopes with at least one turning point in between;

2) The signal part under consideration deviates at least $30\mu V$ from the reference level for a duration of at least 6ms;

3) The minimum observable duration of wave under consideration is 12ms and amplitude \ge 30µV.

3.4 Calculation after intervals determination

The following parameters are determined according to the requirement of *IEC60601-2-51:2003 Medical electrical equipment - Part 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiographs, Annex GG Definitions and rules for the measurement of ELECTROCARDIOGRAMS.*

No.	Parameter	Calculation
1	HR	$60 / RR^{\odot}$
2	PR-interval	Qs [©] - Ps [®]
3	P-duration	$Pe^{(4)} - Ps^{(3)}$
4	QRS-duration	Se ^⑤ − Qs [∅]
5	T-duration	$\mathrm{Te}^{\overline{\mathrm{C}}}$ - $\mathrm{Ts}^{\mathrm{@}}$

6	QT	$Te^{\overline{C}} - Qs^{@}$					
7	QTc	$rac{QT}{\sqrt{RR}}$ \odot					
8	P/QRS/T electric axis	Electric axis formula: $\frac{\arctan(2.0 \times (S_{III} + S_{I}), S_{I} \times \sqrt{3}) \times 180}{PI}$ (8) P electric axis: S_{III} : voltage sum from the beginning point to the end point of P-wave on lead III S_{I} : voltage sum from the beginning point to the end point of P-wave on lead I QRS electric axis: S_{III} : voltage sum from the beginning point to the end point of QRS-complex on lead III S_{I} : voltage sum from the beginning point to the end point of QRS-complex on lead III T electric axis: S_{III} : voltage sum from the beginning point to the end point of QRS-complex on lead I T electric axis: S_{III} : voltage sum from the beginning point to the end point of T-wave on lead III S_{I} : voltage sum from the beginning point to the end point of T-wave on lead III					
9	R(V5)	Height (voltage value) of R-wave on lead V5					
10	S(V1)	Height (voltage value) of S-wave on lead V1					

ANote:

- 1 RR: RR-interval
- 2 Qs: beginning of the Q-wave
- ③ Ps: beginning of the P-wave
- 4 Pe: end of the P-wave
- ⑤ Se: end of the S-wave
- ⁽⁶⁾ Ts: beginning of the T-wave
- 1 Te: end of the T-wave
- (8) PI: 3.1415926

3.5 Interpretations judgment based on parameters

No.	Item	Rule of interpretation
1	No abnormal	No any abnormal are detected

2	Sinus mode Bradycardia	Sinus P-wave, PR-interval between
		110ms-210ms, HR≤*/min, general *=50
3	Sinus mode Tachycardia	Sinus P-wave, PR-interval between
		110ms-210ms, HR \geq */min, general *=100
		P-wave of leads I, II, aVL shall meet the
4	Left atrium Hypertrophy	conditions: width increase of P-wave≥110ms,
		or P-wave displays in double-peak type,
		value of peak to peak ≥40ms
5	Right atrium Hypertrophy	For leads I, II, aVF, amplitude of P-wave
		$\geq 0.25 \text{mV}$, or P-wave is sharp
6	Dual atrium Hypertrophy	
0		
7	QRS low voltage	Voltage of I-aVF limb leads < 0.5 mV,
,	QKS low voltage	e Voltage of I-aVF limb leads <0.5m²
8	Cardiac electric axis normal	QRS-axis between 30 to 90 degree
9	Left axis deviation	QRS-axis between -90 to-30 degree
10	Right axis deviation	QRS-axis between 120 to 180 degree
11	Completeness Right Bundle branch	QRS-duration>120ms, R-wave of lead
11	block	V1 or aVR is wide (width of R-wave>80ms)
	Completeness Left Bundle branch	QRS-duration>120ms, R-wave of lead
12	block	V5 or V6 is wide
	No Completeness Right Bundle	ORS-duration<120ms, R-wave of lead
13	branch block	V1 or aVR is wide (width of R-wave>80ms)
-	No Completeness Left Bundle	QRS-duration<120ms, R-wave of lead V15
14	branch block	or V6 is wide (width of R-wave>80ms)
15	V1 shows RSR' type	QRS-complex of lead V1 is RSR' type
	vi slows Rok type	QRS-duration<110ms, QRS-axis <-30
		degree, lead I and lead aVL are qR type, and
16	Left anterior fascicular block	Q-wave duration<20ms, lead II, III and aVF
		are rS type.
		QRS-duration<110ms, QRS-axis >90
		degree, lead I and lead aVL are rS type, lead
17	Left posterior fascicular block	II, III and aVF are qR type, and Q-wave of
		lead II and III <20ms.
		R amplitude of lead I >1.5mV, R
		amplitude of lead V5 >2.5mV, R amplitude
		of lead aVL >1.2mV, R amplitude of lead
18	Left ventricular hypertrophy	· •
		aVF >2mV, R amplitude of lead V5 minus S
		amplitude of lead V1 >4mV (male) or $3.5mV$
<u> </u>		(female).
		R amplitude of lead aVR >0.5mV, R
19	Right ventricular hypertrophy	amplitude of lead V1 >1mV, R amplitude of
	6	lead V1 minus S amplitude of lead
		V5 >1.2mV, R amplitude of lead V1 is larger

		than S amplitude, R amplitude of lead V5 is
		smaller than S amplitude.
20	I atrioventricular block	PQ interval >210ms
21	Early anteroseptal MI	Early myocardial infarction change of leads V1, V2, V3, no change of leads V4, V5.
22	Possible acute forepart anteroseptal MI	Acute myocardial infarction change of leads V1, V2, V3, no change of leads V4, V5.
23	Old anteroseptal MI	Old myocardial infarction change of leads V1, V2, V3, no change of leads V4, V5.
24	Early anterior MI	Early myocardial infarction change of leads V3, V4, V5, no change of leads V1, V2, V6.
25	Possible acute anterior MI	Acute myocardial infarction change of leads V3, V4, V5, no change of leads V1, V2, V6.
26	Old anterior MI	Old myocardial infarction change of leads V3, V4, V5, no change of leads V1, V2, V6.
27	Early extensive anterior MI	Early myocardial infarction change of leads V1, V2, V3, V4, V5.
28	Possible acute extensive anterior MI	Acute myocardial infarction change of leads V1, V2, V3, V4, V5.
29	Old extensive anterior MI	Old myocardial infarction change of leads V1, V2, V3, V4, V5.
30	Early apical MI	Early myocardial infarction change of leads V4, V5, no change of leads V1, V2, V3.
31	Acute apical MI	Acute myocardial infarction change of leads V4, V5, no change of leads V1, V2, V3.
32	Old apical MI	Old myocardial infarction change of leads V4, V5, no change of leads V1, V2, V3.
33	Early anterolateral MI	Early myocardial infarction change of leads I, aVL, V4, V5, V6
34	Possible acute anterolateral MI	Acute myocardial infarction change of leads I, aVL, V4, V5, V6.
35	Old anterolateral MI	Old myocardial infarction change of leads I, aVL, V4, V5, V6
36	Early high lateral MI	Early myocardial infarction change of leads I, aVL, no change of leads II, III, aVF, V4, V5, V6.
37	Possible acute high lateral MI	Acute myocardial infarction change of leads I, aVL, no change of leads II, III, aVF, V4, V5, V6.
38	Old high lateral MI	Old myocardial infarction change of leads I, aVL, no change of leads II, III, aVF, V4, V5, V6.
39	Early inferior MI	Early myocardial infarction change of leads II, III, aVF, no change of leads I, aVL.

40	Possible acute inferior MI	Acute myocardial infarction change of leads II, III, aVF, no change of leads I, aVL.
41	Old inferior MI	Old myocardial infarction change of leads II, III, aVF, no change of leads I, aVL.
42	Early inferolateral MI	Early myocardial infarction change of leads I, II, III, aVL, aVF.
43	Possible acute inferolateral MI	Acute myocardial infarction change of leads I, II, III, aVL, aVF.
44	Old inferolateral MI	Old myocardial infarction change of leads I, II, III, aVL, aVF.
45	ST depression, mild anteroseptal myocardial ischemia	Mild ST-segment depression of leads V1, V2, V3, and no change of leads V4, V5.
46	ST depression, mild anterior myocardial ischemia	Mild ST-segment depression of leads V3, V4, V5, and no change of leads V1, V2, V6.
47	ST depression, mild extensive anterior myocardial ischemia	Mild ST-segment depression of leads V1, V2, V3, V4, V5.
48	ST depression, mild apical myocardial ischemia	Mild ST-segment depression of leads V4, V5, and no change of leads V1, V2, V3.
49	ST depression, mild anterolateral myocardial ischemia	Mild ST-segment depression of leads I, aVL, V4, V5, V6.
50	ST depression, mild high lateral myocardial ischemia	Mild ST-segment depression of leads I, aVL, and no change of leads II, III, aVF, V4, V5, V6.
51	ST depression, mild inferior myocardial ischemia	Mild ST-segment depression of leads II, III, aVF, and no change of leads I, aVL.
52	ST depression, mild inferolateral myocardial ischemia	Mild ST-segment depression of leads I, II, III, aVL, aVF.
53	ST depression, anteroseptal myocardial ischemia	Severe ST-segment depression of leads V1, V2, V3, and no change of leads V4, V5.
54	ST depression, anterior myocardial ischemia	Severe ST-segment depression of leads V3, V4, V5, and no change of leads V1, V2, V6.
55	ST depression, extensive anterior myocardial ischemia	Severe ST-segment depression of leads V1, V2, V3, V4, V5.
56	ST depression, apical myocardial ischemia	Severe ST-segment depression of leads V4, V5, and no change of leads V1, V2, V3.
57	ST depression, anterolateral myocardial ischemia	Severe ST-segment depression of leads I, aVL, V4, V5, V6.
58	ST depression, high lateral myocardial ischemia	Severe ST-segment depression of leads I, aVL, and no change of leads II, III, aVF, V4, V5, V6.
59	ST depression, inferior myocardial ischemia	Severe ST-segment depression of leads II, III, aVF, and no change of leads I, aVL.
60	ST depression, inferolateral myocardial ischemia	Severe ST-segment depression of leads I, II, III, aVL, aVF.

ANote:

Early myocardial infarction: normal Q-wave, ST elevation or ST slope elevation Acute myocardial infarction: abnormal Q-wave, ST elevation or ST slope elevation Old myocardial infarction: abnormal Q-wave, no ST elevation. Abnormal Q-wave:

For leads I, II, III, avR, avL, avF, V3, V4, V5, V6, voltage of Q-wave <-0.3mV, or 4 times of negative wave of Q-wave> voltage of R-wave and R'-wave, and/or Q-duration>40ms.

For leads V1, V2, voltage of Q-wave <-0.08mV and Q-duration>10ms.

ST elevation:

For leads I, II, III, avR, avL, avF, V4, V5, V6, the voltage of ST segment at 60ms point >0.1mV, and for leads V1, V2, V3, the voltage at 60ms point >0.3mV.

ST slope elevation:

Voltage of ST segment at 20ms point>=voltage of J point, voltage at 40ms point >= the one at 20ms, voltage at 60ms point >= the one at 40ms, with change of ST elevation.

4. Data sources and data preprocessing

4.1 Data sources

According to the requirement of *IEC60601-2-51:2003 Medical electrical equipment - Part* 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiograph, the CSE measurement database, CSE diagnostic database, CTS calibration database and customized data shall be used to evaluate the function of automated measurements and automated interpretations.

Verification	Database	Database items
		CAL05000 CAL10000 CAL15000 CAL20000
	CTS database	CAL20002 CAL20100 CAL20110 CAL20160
Automated	C15 Ualabase	CAL20200 CAL20210 CAL20260 CAL20500
measurement		CAL30000 ANE20000 ANE20001 ANE20002
	CSE measurement	MA 0001~MA0125
	database	MA_0001 - MA0123
Automated	CSE diagnostic database	D_0001~D_1220
interpretation	Customized data	000001~000549

4.2 CTS introduction

The CTS computerized ECG conformance testing project was launched in 1989 by the European Union. This project laid the foundation for computerized ECG conformance testing service. Currently, about 20 types of waveform have been designed derived from the test signals having an infinite length, these signals are part of the CTS-ECG test database, and have proven their effectiveness in a series of official tests. According to the requirement of *IEC60601-2-51:2003 Medical electrical equipment - Part 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiograph Clause 50.101.1*, 13 data (CAL05000, CAL10000, CAL15000, CAL20000, CAL20002, CAL20100, CAL20110, CAL20160, CAL20200, CAL20210, CAL20260, CAL20500, CAL30000) are used in the automated parameters verification for this test.

4.3 CSE introduction

The EU CSE (Common Standards for Quantitative Electrocardiography) ECG database contains 3-lead measurement database of collection1 and collection2, 12-lead measurement database of collection3 and collection4, and a diagnostic database of collection5. In which, the 12-lead measurement database contains 250 groups of interference data; Diagnostic database contains 1220 cases of short-term ECG recording. The primary development purpose of using 12-lead or 15-lead is to evaluate the performance of the automatic ECG analyzer. In addition to the normal data, the database also includes clinically confirmed ECGs of variety cases, such as left ventricular hypertrophy, right ventricular hypertrophy, every part of myocardial infarction and ventricular hypertrophy accompanying myocardial infarction. The database has made a great contribution to the study of electrocardiology, which is, the CSE group published a report on the recommended standard for general ECG measurements based on the investigation and study of the database, which has been widely recognized by the world.

Item	Number
Normal	382
Left ventricular hypertrophy	183
Right ventricular hypertrophy	55
Biventricular hypertrophy	53
Anterior myocardial infarction	170
Inferior myocardial infarction	273
Complex myocardial infraction	104
Synthetical accuracy	1220

CSE database diagnostic items:

4.4 Customized data

4.4.1 Data description

Customized	Description
data	
Total recording	549
number	
Race	Yellow race
<i>a i</i>	Aged from 17 to 87, average age 57.23, standard deviation 21.32;
Coverage of	326 male, average age 55.54, standard deviation 19.81;
age, gender	223 female, average age 59.70, standard deviation 22.63.
	12-lead ECG data (I, II, III, AVR, AVL, AVF, V1, V2, V3, V4, V5, V6),
Sampling data	sampling frequency of each channel: 1kHz, amplitude quantization:
	2.4µV/LSB.
	The interpretation conclusion of customized data is determined by the
	physician diagnostic results of cardiac catheterization and ultrasonic
	examination, and the ECG judgment result in physical examination, the
Remark	details as blow:
	1) Normal ECG
	Determined by the diagnostic result that judged as normal in cardiac
	catheterization and ultrasonic examination, and the result that judged as

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ANote:

The heart abnormalities such as posterior myocardial ischemia, early posterior MI and old posterior MI are not included in the database. These abnormalities and other heart disorders not contained in above sheet won't be regarded as the judgment object for the verification of automated interpretation accuracy.

4.6 Data preprocessing

4.6.1 CTS preprocessing

The 16 cases (CAL05000, CAL10000, CAL15000, CAL20000, CAL20002, CAL20100, CAL20110, CAL20160, CAL20200, CAL20210, CAL20260, CAL20500, CAL30000, ANE20000, ANE20001, ANE20002) from CTS-ECG shall be processed for voltage conversion and frequency conversion for resampling as the applicable format in the system. Then cases will be imported to the device. After that, the verification of automated measurement parameters will be carried on.

4.6.2 CSE preprocessing

The cases (MA_0001~MA0125, D_0001~D_1220) from the CSE shall be processed for voltage conversion and frequency conversion for resampling as the applicable format in the system. Then cases will be imported to the device. After that, the case of MA_0001~MA0125 shall be used for the following verification of automated measurement parameters, and the case of D_0001~D_1220 shall be used for the following verification of automated interpretation.

4.6.3 Customized data preprocessing

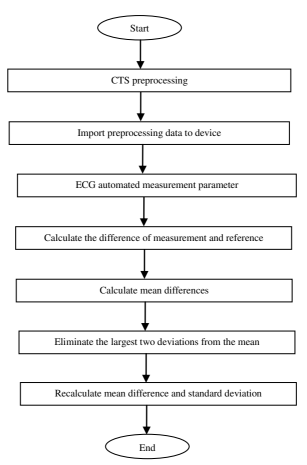
The customized initial case files shall be processed for voltage conversion and frequency conversion for resampling as the applicable format in the system. Then cases will be imported to the device. After that, the verification of automated interpretation will be carried on.

5. Process and Result of Verification

5.1 Verification of measurement function

5.1.1 Verification and Process for CTS measurement database

The cases (CAL05000, CAL10000, CAL15000, CAL20000, CAL20100, CAL20110, CAL20160, CAL20200, CAL20210, CAL20260, CAL20500, CAL20500, ANE20000, ANE20001, ANE20002) imported to the device shall be used to verify the automated measurement parameters.



5.1.2 Verification and Process for CSE measurement database

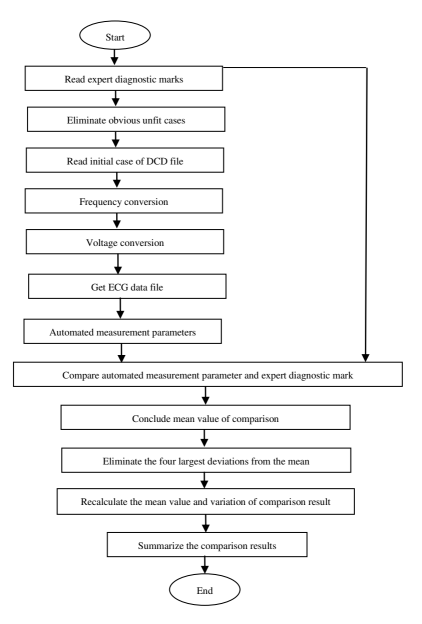
Import the converted case files into the device, add appropriate database records, then waveform for all case files can be reviewed in the device, therefore the automated measurement parameters can be obtained.

Eliminate the cases existing obvious error for the diagnostic parameters (P-wave location is wrong) from the CSE database.

Make a comparison between the ECG analytical parameters (the beginning/end of P-wave, QRS-complex and T-wave) and the diagnostic parameters (the beginning/end of P-wave, QRS-complex and T-wave) provided by CSE database. Draw the two groups of waveform and mark the location of the beginning/end of P-wave, QRS-complex and T-wave corresponding to each case. The picture provides a visualized comparison, so the mean and standard deviation of the differences can be calculated. According to the requirement of IEC60601-2-51:2003 Medical electrical equipment - Part 2-51: Particular requirements for safety, including essential performance, of recording and analysing single channel and multichannel electrocardiograph, the

four largest deviations from the mean shall be eliminated before recalculation of mean and standard deviation of the differences.

Flow diagram of CSE measurement database verification process



5.1.3 Verification results

5.1.3.1 Accuracy of amplitude measurements

Calibration and analytical ECGs shall be used to measure the amplitude value, the summary as follows:

Amplitude	Mean difference (uV)	Standard deviation (uV)		
P-wave	-1.70	5.72		
Q-wave	7.51	18.07		
R-wave	-18.05	21.70		
S-wave	7.77	18.58		
ST-segment	0.15	4.24		
T-wave	-5.81	8.03		

Mote: In amplitude measurement, for large-amplitude ECG, such as CAL30000, it is

necessary to adjust to 0.5 times the gain before testing.

5.1.3.2 Accuracy of absolute interval and wave duration measurements

Calibration and analytical ECGs shall be used to measure the global interval and wave duration (including Q-wave ,R-wave ,S-wave), the summary as follows:

Interval&Duration	Mean difference (ms)	Standard deviation (ms)
P-duration	-5.70	1.88
PQ-interval	-2.58	1.94
QRS-duration	-0.23	3.26
QT-interval	-6.70	4.37

5.1.3.3 Accuracy of interval measurements on biological ECGs

CSE database shall be used to evaluate the accuracy of interval measurements on biological ECGs, the summary as follows:

Interval&Duration	Mean difference (ms)	Standard deviation (ms)
P-duration	0.99	13.46
PR-interval	3.65	9.68
QRS-duration	-1.69	6.11
QT-interval	-2.32	20.69

5.1.3.4 Stability of measurements against NOISE

The test is carrying on according to MA-series data (008, 011, 013, 014, 015, 021, 026, 027, 042, 061) in CSE database.

Global measurement	Type of added	Disclosed differences		
parameters	NOISE	Mean (ms)	Standard deviation (ms)	
P-duration	High frequency	-5.65	12.33	
P-duration	Line frequency	-0.25.	12.71	
P-duration	Base-line	-4.90	33.15	
QRS-duration	High frequency	-0.95	5.13	
QRS-duration	Line frequency	1.35	4.71	
QRS-duration	Base-line	-1.55	7.68	
QT-interval	High frequency	-14.55	6.51	

QT-interval	Line frequency	-8.55	20.73
QT-interval	Base-line	36.20	64.47

The biological ECGs are fed into the device in form of digital signals, then the measurement value can be obtained by calculation.

Test condition:

a) without NOISE

b)with 25uV high frequency

c) with 50uV peak to valley 50Hz/60Hz sinusoidal line frequency NOISE

d) with 1mV peak to valley 0.3Hz sinusoidal base-line NOISE

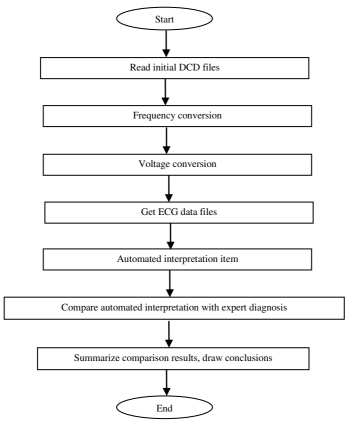
For each NOISE level above, the differences of measurements between the NOISE-free ECGs and the ECGs with NOISE shall be determined. The two largest deviations from the mean

shall be estimated before calculation of mean and standard deviation of differences.

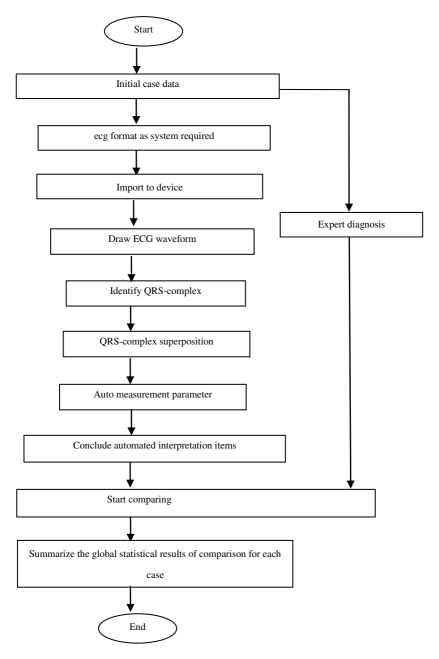
5.2 Verification of interpretation function

5.2.1 Verification process

5.2.1.1 CSE diagnostic database



5.2.1.2 Customized database



5.2.2 Verification results

No.	Item	ECGs number	Sensitivit y %	Specific ity %	Positive predictive value %
1	No abnormal	585	92.01	79.16	97.38
2	Sinus mode Bradycardia	191	96.68	99.73	98.64
3	Sinus mode Tachycardia	78	97.44	96.49	96.90
4	Left atrium Hypertrophy	51	51.09	99.89	81.82
5	Right atrium Hypertrophy	43	42.64	99.66	50.00
6	Dual atrium Hypertrophy	22	93.58	99.14	60.19
7	QRS low voltage	5	96.37	99.36	63.25
8	Cardiac electric axis normal	733	98.36	89.13	98.79
9	Left axis deviation	168	98.65	89.40	98.18
10	Right axis deviation	107	98.23	88.99	94.90
11	Completeness Right Bundle branch block	28	97.00	89.50	95.45
12	Completeness Left Bundle branch block	32	97.73	89.65	91.43
13	No Completeness Right Bundle branch block	41	96.86	89.83	82.35
14	No Completeness Left Bundle branch block	47	94.68	89.83	89.66
15	V1 shows RSR' type	13	90.32	91.14	65.12
16	Left anterior fascicular block	26	91.43	93.25	71.11
17	Left posterior fascicular block	18	89.29	97.37	52.63
18	Left ventricular hypertrophy	236	41.37	92.65	70.36
19	Right ventricular hypertrophy	108	39.75	93.47	65.39
20	I atrioventricular block	13	94.58	91.67	80.64
21	Early anteroseptal MI	10	83.33	99.94	90.91
22	Possible acute forepart anteroseptal MI	27	16.67	98.73	91.89
23	Old anteroseptal MI	26	92.00	98.90	86.47
24	Early anterior MI	77	93.90	88.22	71.96
25	Possible acute anterior MI	10	80.00	99.72	44.44
26	Old anterior MI	13	24.00	99.66	50.00
27	Early extensive anterior MI	24	79.67	99.43	41.18
28	Possible acute extensive anterior MI	16	81.82	99.66	75.00
29	Old extensive anterior MI	30	90.91	88.05	37.04
30	Early apical MI	15	88.32	87.21	88.54
31	Acute apical MI	21	78.12	78.66	53.85
32	Old apical MI	19	79.63	89.94	80.00
33	Early anterolateral MI	36	77.51	79.94	83.33

	1		1	1	
34	Possible acute anterolateral MI	9	28.57	99.77	33.33
35	Old anterolateral MI	14	70.00	93.60	50.00
36	Early high lateral MI	16	79.65	95.78	80.42
37	Possible acute high lateral MI	8	81.60	99.94	85.71
38	Old high lateral MI	23	81.82	99.66	60.00
39	Early inferior MI	31	88.89	95.00	40.00
40	Possible acute inferior MI	11	76.00	99.60	61.11
41	Old inferior MI	101	96.07	99.24	93.44
42	Early inferolateral MI	73	98.77	96.82	75.94
43	Possible acute inferolateral MI	29	11.11	99.94	50.00
44	Old inferolateral MI	28	84.62	99.83	78.57
45	ST depression, mild anteroseptal myocardial ischemia	7	75.36	99.55	46.67
46	ST depression, mild anterior myocardial ischemia	5	81.24	99.94	33.33
47	ST depression, mild extensive anterior myocardial ischemia	13	79.83	99.13	53.59
48	ST depression, mild apical myocardial ischemia	17	76.97	99.14	43.13
49	ST depression, mild anterolateral myocardial ischemia	25	77.54	99.08	37.64
50	ST depression, mild high lateral myocardial ischemia	21	80.64	99.14	47.39
51	ST depression, mild inferior myocardial ischemia	12	79.73	99.60	55.16
52	ST depression, mild inferolateral myocardial ischemia	20	80.59	99.26	50.61
53	ST depression, anteroseptal myocardial ischemia	4	85.41	99.72	44.44
54	ST depression, anterior myocardial ischemia	12	87.66	98.58	34.85
55	ST depression, extensive anterior myocardial ischemia	7	84.78	98.04	67.75
56	ST depression, apical myocardial ischemia	18	79.95	99.14	55.12
57	ST depression, anterolateral myocardial ischemia	13	87.42	98.97	59.09
58	ST depression, high lateral myocardial ischemia	16	90.06	99.31	57.14
59	ST depression, inferior myocardial ischemia	12	89.88	99.13	40.08
60	ST depression, inferolateral myocardial ischemia	6	91.39	99.16	50.47

Sensitivity: probability that a "True sample" would be determined as certain "Item" by

automated interpretation function;

Specificity: probability that a "True unfit sample" would be determined as certain "Unfit item" by automated interpretation function;

Positive predictive value: probability that a determined "Unfit item" is a "True unfit item".

6. Accuracy of rhythm diagnosis

6.1 ECG database used for rhythm diagnosis

The ECG database that used for testing the accuracy of rhythm diagnosis contains 3000 cases of 12-lead ECG, each case length is 10s. The data is measured by 12-lead ECG device of our company. The true value of data is judged by a heart expert with more than 10 years of work experience based on those 12-lead ECG waveform.

The number of cases with the following diagnosis types (the case diagnosis may contain one or more types) is shown as below:

Rhythm type	Number
Sinus rhythm	2003
Sinus Tachycardia	313
Sinus Bradycardia	338
Arrhythmia	112
Ventricular premature beat	230
Ventricular premature beat (double)	16
Ventricular premature beat (bigeminy)	140
Ventricular premature beat (trigeminy)	147
Ventricular tachycardia	42
Atrial fibrillation	232

Other rhythm types that not included in the database: atrial flutter, ventricular fibrillation, supraventricular rhythm, junctional rhythm, pacemaker rhythm, II °/III° atrioventricular block, arrest and other ECG abnormal.

The statistical information of ECG database that used for testing the accuracy of rhythm diagnosis is shown as below:

		Total					Male			Female					
	Yo	Ol	Av	S	Т	You	Ol	Av	S	Т	Yo	Ol	Av	S	Т
	ung	de	era	D	ot	nges	de	era	D	ot	ung	de	era	D	ot
	est	st	ge		al	t	st	ge		al	est	st	ge		al
Т				1	3				1					1	1
	10	02	48.	1	0	10	02	50.	7	14	10	02	47.	8	5
ot	12	93	6	7. 9	0	12	93	0		95	12	92	3		0
al				9	0				7					0	5

6.2 Verification results of accuracy of rhythm diagnosis

The ECG obtained from the ECG database for rhythm diagnosis is input to the electrocardiograph for testing in form of digital signals. The rhythm results analyzed by electrocardiograph are compared with the true rhythm results of ECG, and the calculated sensitivity, specificity and positive predictive value are shown as below:

Rhythm type	ECGs number	Sensitivity %	Specificity %	Positive predictive value %
Sinus rhythm	3000	83.82	97.79	98.71
Arrhythmia	3000	75.89	98.86	72.03
Tachycardia	3000	96.81	95.27	70.47
Bradycardia	3000	99.11	99.44	95.71
Ventricular tachycardia	3000	83.33	99.73	81.4
Ventricular premature beat	3000	81.3	98.3	79.91
Ventricular premature beat (double)	3000	87.5	99.87	77.78
Ventricular premature beat (bigeminy)	3000	93.57	99.55	90.97
Ventricular premature beat (trigeminy)	3000	88.44	99.82	96.3
Atrial fibrillation	3000	50.86	98.55	74.68

ANote:

Sensitivity: probability that a "True sample" would be determined as certain "Rhythm type" by rhythm diagnosis function;

Specificity: probability that a "True unfit sample" would be determined as certain "Unfit rhythm type" by rhythm diagnosis function;

Positive predictive value: probability that a determined "Unfit rhythm type" is a "True unfit rhythm type".

Appendix II EMC Guidance and Manufacturer Declaration

Table 1:

Table 1:			
Guidance and manufa	cturer	's declaration -electrom	agnetic emission
This machine is intended for use	in the	e electromagnetic envir	conment specified below. The
purchaser or the user of the device	should	assure that it is used in	such environment.
Emission test		Compliance	
RF emissions CISPR 11		Group 1	
RF emissions CISPR 11		Class A	
Harmonic emissions IEC 61000-3-2		Class A	
Voltage fluctuations/flicker emission IEC 61000-3-3	ons	Not applicable	
Table 2:			
Guidance and manufa	cturer	's declaration-electroma	agnetic immunity
This machine is intended for use purchaser or the user of this machin		e	1
Immunity test	IEC6 test	0601 level	Compliance level
Electrostatic discharge (ESD)	±8kV	⁷ contact	±8kV contact
IEC 61000-4-2	± 15	kV air	±15kV air
Electrical fast transient/burst IEC 61000-4-4	lines	⁷ for power supplyV for input/output line	±2kV for power supply lines Not Applicable
Surge	±1 kV	V lines to lines	±1 kV lines to lines
IEC 61000-4-5	±2 kV	V lines to earth	±2 kV lines to earth
		UT(>95%dip in UT) 5 cycle	<5%UT(>95%dip in UT) for 0.5 cycle
Voltage dips, short interruptions and voltage vatiations on power supply input lines IEC 61000-4-11	40% 5 cyc	UT(60%dip in UT) for le UT(30%dip in UT) for	40% UT(60%dip in UT) for 5 cycle 70%UT(30%dip in UT) for 25 cycle
110 01000-4-11	-	UT(>95%dip in UT)	<5%UT(>95%dip in UT) for 5 sec
Power frequency (50 / 60Hz) magnetic field IEC 61000-4-8	30 A		30A/m

Table 3:

Guidance and	d manufacturer	's declaration	- electromagne	tic immunity

This machine is intended for use in the electromagnetic environment specified below. The customer the user of this machine should assure that it is used in such environment.

Immunity test	IEC 60601 test level	Compliance level		
Conducted RF IEC61000-4-6	3 V 0,15 MHz – 80 MHz 6 V in ISM bands between 0,15 MHz and 80 MHz	3 V 0,15 MHz – 80 MHz 6 V in ISM bands between 0,15 MHz and 80 MHz		
Radiated RF IEC61000-4-3	3 V/m 80 MHz- 2.7 GHz	3 V/m80 MHz- 2.7 GHz		

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which this machine is used exceeds the applicable RF compliance level above, this machine should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating this machine.

Table 4:

Guidance and manufacturer's declaration - electromagnetic Immunity							
The [Code SI] is intended for use in the electromagnetic environment specified below. The customer							
or the user of the [Code SI] should assure that it is used in such an environment							
	-						

Radiated RF IEC6100	Test Frequency (MHz)	Band a) (MHz)	Service a)	Modulation b)	Modulation b) (W)	Distance (m)	IMMUNITY TEST LEVEL (V/m)
0-4-3 (Test specificat ions for	385	380 -390	TETRA 400	Pulse modulation b) 18 Hz	1,8	0,3	27
ENCLOS URE PORT IMMUNI	450	380 -390	GMRS 460, FRS 460	FM c) ± 5 kHz deviation 1 kHz sine	2	0,3	28
TY to RF wireless communi	710 745 780	704 – 787	LTE Band 13, 17	Pulse modulation b) 217 Hz	0,2	0,3	9

			~~~				
cations	810		GSM				
equipmen			800/900,			0,3	28
t)	870		TETRA	<b>D</b> 1	2		
	070		800,	Pulse			
		800 -	iDEN	modulation			
		960	820,	b)			
	930		CDMA	18 Hz			
	950		850,				
			LTE				
		-	Band 5				
	1720		GSM				
	1845		1800;			0,3	28
	1045		CDMA				
		1 700 – 1 990	1900;		2		
			GSM				
			1900;				
	1970		DECT;				
			LTE				
			Band 1, 3,				
			4, 25;				
			UMTS				
			Bluetooth				
			,		2	0,3	28
		2 400 - $802.2 570 b/g/r$	WLAN,	Pulse modulation b) 217 Hz			
			802.11				
	2450 2 570		b/g/n,				
			RFID				
		5 100 – 5 800	2450,				
			LTE				
			Band 7				
	5240		WLAN	Pulse	0,2	0,3	9
	5500		802.11	modulation b)			
			a/n				
	5785			217 Hz	<u> </u>		
		achieve th	e IMMUNI	TY TEST LEV	EL, the distand	ce between	the transmitting
antenna and the							

ME EQUIPMENT or ME SYSTEM may be reduced to 1 m. The 1 m test distance is permitted by IEC 61000-4-3.

a) For some services, only the uplink frequencies are included.

b) The carrier shall be modulated using a 50 % duty cycle square wave signal.

c) As an alternative to FM modulation, 50 % pulse modulation at 18 Hz may be used because while it does not represent actual modulation, it would be worst case.

The MANUFACTURER should consider reducing the minimum separation distance, based on RISK MANAGEMENT, and using higher IMMUNITY TEST LEVELS that are appropriate for the reduced minimum separation distance. Minimum separation distances for higher IMMUNITY TEST LEVELS shall be calculated using the following equation:

$$E = \frac{6}{d}\sqrt{P}$$

Where P is the maximum power in W, d is the minimum separation distance in m, and E is the IMMUNITY TEST LEVEL in V/m.

**M**Warning

- Don't near active HF SURGICAL EQUIPMENT and the RF shielded room of an ME SYSTEM for magnetic resonance imaging, where the intensity of EM DISTURBANCES is high.
- 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.
- 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."
- 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 device including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result.
- Active medical devices are subject to special EMC precautions and they must be installed and used in accordance with these guidelines.

**Mote:** 

- The EMISSIONS characteristics of this equipment make it suitable for use in industrial areas and hospitals (CISPR 11 class A). If it is used in a residential environment (for which CISPR 11 class B is normally required) this equipment might not offer adequate protection to radio-frequency communication services. The user might need to take mitigation measures, such as relocating or re-orienting the equipment.
- When the device is disturbed, the data measured may fluctuate, please measure repeatedly or in another environment to ensure its accuracy.

#### § 15.19 Labeling requirements.

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.

#### § 15.21 Information to user.

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

### § 15.105 Information to the user.

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 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 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.

-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.

-Consult the dealer or an experienced radio/TV technician for help.

This device need additional AC Power line and fixed at a desk or a cabinet, the wifi transmitter will be keep a 20cm distance to the human body.

This equipment complies with FCC radiation exposure limits set forth for an uncontrolled environment. This equipment should be installed and operated with minimum distance 20cm between the radiator & your body.