

EDAN Agile PLM Electronic Signature Information

--Signatures related to this document and performed in EDAN Agile PLM.

文件名称(Document Name) : F9说明书_英文

文件编号(Number) : 01.54.107779

版本(Version) : 2.9

产品型号(Product Model) : F9;F9 Express

项目编码(Project Code) : 2040A000

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F9, F9 Express

Fetal & Maternal Monitor

User Manual

CE₀₁₂₃



About this Manual

P/N: 01.54.107779

Version 2.9

MPN: 01.54.107779029

Release Date: Sep. 2018

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Statement

This manual will help you understand the operation and maintenance of the product better. It is reminded that the product shall be used strictly complying with this manual. User's operation failing to comply with this manual may result in malfunction or accident for which EDAN INSTRUMENTS, INC. (hereinafter called EDAN) can not be held liable.

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Assembly operations, extensions, re-adjustments, modifications or repairs are carried out by persons authorized by EDAN, and

The electrical installation of the relevant room complies with national standards, and

The instrument is used in accordance with the instructions for use.

Terms Used in this Manual

This guide is designed to give key concepts on safety precautions.

WARNING

A **WARNING** label advises against certain actions or situations that could result in personal injury or death.

CAUTION

A **CAUTION** label advises against actions or situations that could damage equipment, produce inaccurate data, or invalidate a procedure.

NOTE

A **NOTE** provides useful information regarding a function or a procedure.

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Chapter 1 Safety Guidance

CAUTION

Federal (U.S.) Law restricts this device to sale by or on the order of a physician.

NOTE:

- 1 In order to ensure the operator and patient's safety, read through this chapter before using this monitor.
- 2 This user manual is written to cover the maximum configuration. Therefore, your model may not have some of the parameters and functions described, depending on what you have ordered.
- 3 The functions frequently used are marked with an asterisk *, for example 4.10 *Reviewing Alarms.

The monitor mainly consists of the main unit, FTS-3 fetal telemetry system (including the base station, wireless US-T transducers, wireless TOCO-T transducer or TOCO-E transducer, DECG cable and MECG cable), built-in recorder, recorder paper, US transducer, TOCO transducer, ECG lead, NIBP cuff, SpO₂ sensor, TEMP sensor, event remote marker, fetal stimulator, IUP cable and catheter and other accessories.

The monitor is intended to monitor FHR, DECG, fetal movement, TOCO, ECG, PR, IUP, NIBP, SpO₂, TEMP.

1.1 Intended Use/Indications for Use

F9 Fetal & Maternal Monitor (hereinafter called F9):

The **F9** Fetal & Maternal Monitor is intended for non-invasive and invasive monitoring of fetus during antepartum examination, labor and delivery. It is intended to be used only by trained and qualified personnel in antepartum examination rooms, labor and delivery rooms.

F9 Fetal & Maternal Monitor provides Non-Stress testing for pregnant women. It can externally monitor the FHRs using ultrasound and uterine activity via a TOCO transducer. Alternatively, it can internally monitor one of the FHRs with DECG and uterine activity with an IUPC.

F9 Express Fetal & Maternal Monitor (hereinafter called F9 Express):

F9 Express Fetal & Maternal Monitor is intended for monitoring physiological parameters of pregnant women during antepartum examination, labor and delivery. It is intended to be used only by trained and qualified personnel in antepartum examination rooms, labor and delivery rooms.

F9 Express Fetal & Maternal Monitor is intended for providing Non-Stress testing or fetal monitoring for pregnant women. In addition, it provides a solution for maternal vital signs

monitoring.

Contraindications:

They are not intended for use in intensive care units, operating rooms or for home use.

1.2 Features

The following table lists the measurements that **F9** and **F9 Express** support.

Measurement	Model	F9	F9 Express
FHR		√	√
TOCO		√	√
FM		√	√
AFM		√	√
DECG		Opt	Opt
IUP		Opt	Opt
MECG		✗	√
NIBP		✗	√
MSpO2		✗	√
TEMP		✗	√
NOTE: √ = Standard Opt = Optional ✗ = Not Available			

For the Essential Performance of **F9** and **F9 Express**, refer to Appendix 1 in details.

1.3 Instruction for Safe Operation

NOTE:

In this manual, **Monitor** refers to both **F9** and **F9 Express**, and is used where the information applies to both models.

- ◆ The monitor is designed to comply with the international safety requirements IEC/EN 60601-1 for medical electrical equipment. It is class I equipment.
- ◆ The monitor operates within specifications at ambient temperatures between +5 °C (+41 °F) and +40 °C (+104 °F). Ambient temperatures that exceed these limits could affect the accuracy of the instrument and cause damage to the modules and circuits. Allow at least 2 inches (5 cm) clearance around the instrument for proper air circulation.
- ◆ You must check that the equipment, cables and transducers do not have visible evidence of damage that may affect patient safety or monitoring capability before each use. If damage is evident, replacement is recommended before use.

- ◆ The monitor must be serviced only by authorized and qualified personnel. The manufacturer does not accept responsibility for safety compliance, reliability and performance if modifications or repairs are carried out by unauthorized personnel. Identical replacement parts must be used.
- ◆ The typical operator's position is in front of the monitor. Please position the device in a location where the operator can easily see the screen and access the operating controls. The protective degree against electric shock of the patient connections is:

Ultrasound (FHR1, FHR2)		
External TOCO		
Fetal Movement Mark (FM)	Type BF	
Fetal Stimulator (FS)		
Intrauterine Pressure (IUP)		
Non-invasive Blood Pressure (NIBP)	Type BF, defibrillation-proof	
Arterial Oxygen Saturation (SpO ₂)		
Direct Electrocardiography (DECG)	Type CF	
Electrocardiography (ECG)	Type CF, defibrillation-proof	
Temperature (TEMP)		

The monitor described in this user manual is not protected against:

- a) The effects of high frequency currents
- b) The interference of electrosurgery equipment

1.4 Ultrasound Safety Guide

◆ Fetal Use

The monitor is designed for continuous fetal heart rate monitoring during pregnancy and labor. Clinical interpretation of fetal heart rate traces can diagnose fetal and/or maternal problems and complications.

◆ Instructions for Use in Minimizing Patient Exposure

The acoustic output of the monitor is internally controlled and can not be varied by the operator in the course of the examination. The duration of exposure is, however, fully under the control of the operator. Mastery of the examination techniques described in the User Manual will facilitate obtaining the maximum amount of diagnostic information with the minimum amount of exposure. The exercising of clinical judgment in the monitoring of low risk patients will avoid unnecessary insonation.

1.5 Safety Precautions

WARNING and **CAUTION** messages must be observed. To avoid the possibility of injury, observe the following precautions during the operation of the instrument.

WARNING

For using safety:

- 1 The monitor or FTS-3 telemetry system (hereinafter called FTS-3) is provided for the use of qualified physicians or personnel professionally trained.
 - 2 The monitor is not intended for use in intensive care units (ICU), operating rooms or for home use.
 - 3 Do not switch on the monitor until all cables have been properly connected and verified.
 - 4 **EXPLOSION HAZARD** - Do not use the monitor or FTS-3 system in the presence of flammable anesthetics or other materials.
 - 5 **SHOCK HAZARD** - The power receptacle must be a three-wire grounded outlet. Never try to adapt the three-prong plug to fit a two-slot outlet. A hospital grade outlet is required. If the outlet has only two slots, make sure that it is replaced with a three-slot grounded outlet before attempting to operate the monitor.
 - 6 Multiple portable socket-outlets shall not be placed on the floor.
 - 7 Additional multiple socket-outlet or extension cord can't be connected to the system.
 - 8 The multiple portable socket-outlet provided with the system shall be only used for supplying power to equipment which is intended to form part of the system. If the electrical device that does not belong to the system plug in the socket, the total power may exceed the maximum load of the separating transformer and cause high temperature and fire. Enclosure leakage current within the system exceeds the standard limit, which may lead an electric risk.
 - 9 **SHOCK HAZARD** - Do not attempt to connect or disconnect a power cord with wet hands. Make certain that your hands are clean and dry before touching a power cord.
 - 10 Do not touch accessible parts of non-medical electrical equipment and the patient simultaneously. Do not touch the signal input or output connector and the patient simultaneously.
 - 11 Accessory equipment connected to the analog and digital interfaces must be certified according to the respective IEC/EN standards (e.g. IEC/EN 60950 for data processing equipment and IEC/EN 60601-1 for medical equipment). Furthermore all configurations shall comply with the valid version of the system standard IEC/EN 60601-1. Anybody who connects additional equipment to the signal input connector or signal output connector to configure a medical system must ensure that the system complies with the requirements of the valid version of the system standard IEC/EN 60601-1. If in doubt, consult our technical service department or your local distributor.
-

WARNING

- 12 Connecting any accessory (such as external printer) or other device (such as the computer) to this monitor makes a medical system. In that case, additional safety measures should be taken during installation of the system, and the system shall provide:
 - a) Within the patient environment, a level of safety comparable to that provided by medical electrical equipment complying with IEC/EN 60601-1, and
 - b) Outside the patient environment, the level of safety appropriate for non-medical electrical equipment complying with other IEC or ISO safety standards.
- 13 Using accessories other than those specified by the manufacturer may result in increased electromagnetic emissions or decreased electromagnetic immunity of the device.
- 14 The device should not be used adjacent to or stacked with other equipment and that if adjacent or stacked use is necessary, the device should be observed to verify normal operation in the configuration in which it will be used.
- 15 All the accessories connected to system must be installed outside the patient vicinity, if they do not meet the requirement of IEC/EN 60601-1.
- 16 Do not exceed the maximum permitted load when using multiple portable socket-outlets to supply the system.
- 17 **SHOCK HAZARD** – Do not connect non-medical electrical equipment, which has been supplied as a part of the system, directly to the wall outlet when the non-medical equipment is intended to be supplied by a multiple portable socket-outlet with an isolation transformer. If multiple instruments are connected to a patient, the sum of the leakage currents may exceed the limits given in the IEC/EN 60601-1 and may pose a safety hazard. Consult your service personnel.
- 18 **SHOCK HAZARD** - Don't connect electrical equipment, which has not been supplied as a part of the system, to the multiple portable socket-outlet supplying the system.
- 19 Do not connect any equipment or accessories that are not approved by the manufacturer or that are not IEC 60601-1 approved to the monitor. The operation or use of non-approved equipment or accessories with the monitor is not tested or supported, and monitor operation and safety are not guaranteed.
- 20 Do not apply this monitor and other ultrasonic equipment simultaneously on a same patient, in case of possible hazard caused by leakage current superposition. Do not apply this monitor simultaneously with other PATIENT-connected equipment, such as, a cardiac pacemaker or other electrical stimulators, on the same patient.
- 21 The monitor can only be used on one patient at a time.
- 22 **SHOCK HAZARD** - Do not remove the top panel cover during operation or while power is connected.
- 23 Equipment and devices that connect to the monitor should form an equipotential body to ensure effective grounding.

WARNING

- 24 Only connect accessories supplied or recommended by the manufacturer to the device.
- 25 The system should be operated by the doctor or under the doctor's instructions.
- 26 Do not apply the monitor during electro-surgery or MRI; otherwise it might result in harming the patient or the operator.
- 27 ECG cables may be damaged when connected to a patient during defibrillation. Check cables for functionality before using them again.
- 28 Only MECG, SpO₂, NIBP and TEMP applied parts of the monitor are defibrillation-proof. When a defibrillator is applied, keep other accessories away from the patient. Otherwise it may result in damaging the monitor or harming the patient.
- 29 After defibrillation, the screen display recovers within 10 seconds if the correct electrodes are used and applied based on the manufacturers' instructions.
- 30 Any non-medical equipment (such as the external printer) is not allowed to be used within the patient vicinity (1.5m/6ft.).
- 31 Make sure that the power is turned off and the power cord is disconnected from the AC socket before connecting or disconnecting equipment. Otherwise, the patient or operator may receive electrical shock or other injury.
- 32 Parts and accessories used must meet the requirements of the applicable IEC 601 series safety standards, and/or the system configuration must meet the requirements of the IEC 60601-1 medical electrical systems standard.
- 33 Do not service or maintain the monitor or any accessory which is in use with the patient.
- 34 Assembly of the monitor and modifications during actual service life shall be evaluated based on the requirements of IEC60601-1.

For proper monitoring:

- 35 Alarms must be set up according to different situations of patients. Make sure that audio sounds can be activated when an alarm occurs.
- 36 Do not perform NIBP measurements on patients with sickle-cell disease or under any condition where the skin is damaged or expected to be damaged.
- 37 Clinical decision making based on the output of the device is left to the discretion of the provider.
- 38 Do not put the sensor on extremities with arterial catheter or venous syringe.
- 39 Do not apply the cuff to a limb that has an intravenous infusion or catheter in place. This could cause tissue damage around the catheter when infusion is slowed or blocked during cuff inflation.
- 40 The disposable accessories are intended to be used only once. Dispose of them properly after use and do not reuse them.
- 41 The spiral electrode and IUP catheter are disposable and should not be reused.

WARNING

- 42 The IUPC is neither intended nor approved for measuring intrauterine pressure extraovularly; attempting to do so may lead to maternal discomfort or injury.

For using the Battery:

- 43 Before using the rechargeable lithium-ion battery (hereinafter called battery), be sure to read the user manual and safety precautions thoroughly.
- 44 Use the battery only in the F9 or F9 Express monitor.
- 45 Do not reverse the battery pole or it will cause explosion.
- 46 Do not unplug the battery when monitoring.
- 47 Do not heat or throw the battery into a fire.
- 48 Before using the battery, make sure to read the user manual and safety precautions thoroughly.
- 49 Do not use or leave battery close to fire or other places where temperatures may be above +60 °C (+140 °F).
- 50 Do not immerse, throw or wet the battery in water/ seawater.
- 51 Do not destroy the battery: Do not pierce battery with a sharp object such as a needle. Do not hit with a hammer, step on or throw or drop to cause strong shock. Do not disassemble or modify the battery.
- 52 Do not short-circuit the battery by connecting the battery cable connector or battery socket with metal objects or solder.
- 53 If the liquid leak from the battery spills onto your skin or clothes, wash well with fresh water immediately.
- 54 If the liquid leak from the battery gets into eyes, do not rub the eyes. Wash them well with clean water and see a doctor immediately.
- 55 Keep the battery away from fire immediately when leakage or foul odor is detected.
- 56 Stop using the battery if abnormal heat, odor, discoloration, deformation or abnormal condition is detected during use, charge, or storage. Keep it away from the monitor.
- 57 Remove the battery and store it at a cool and dry environment if the monitor is not used for a long time.
- 58 Unplug the monitor before installing and removing the battery.
- 59 Do not connect the battery directly to an electric outlet or cigarette lighter charger.
- 60 Batteries have life cycles. If the time that the monitor using battery becomes much shorter than usual, the battery life is at an end. Replace the battery with a new one of the same specification as the one provided or recommended by the manufacturer.
- 61 If the battery is stored alone and not used for a long time, we recommend that the battery should be charged at least once every 6 months to prevent overdischarge.
- 62 Only rechargeable button cell supplied or recommended by EDAN can be used.

WARNING

For network safety:

63 Given the vulnerabilities of WEP, it is advised to use WAP and WAP2. WEP should not be used as it is not secure.

64 Data in transit is no encrypted. Local Area Network isolation or other protections are suggested to prevent unintended exposure of PHI.

In addition, when you use the FTS-3 fetal telemetry system, please pay attention to the warnings as follows:

65 The system should be operated by the doctor or under the doctor's instructions.

66 SHOCK HAZARD – The base station and transducers for one patient must be supplied by the same power and do not change the power supply.

67 Please arrange a function test periodically for the system.

68 Do not move the system when it is powered on and do not soak it in any liquid.

69 Please check the transducer, cable and base station periodically. If the transducers are damaged, do not use them in water and repair them in time.

70 The battery in the wireless transducer should be replaced by the serviceman authorized by EDAN.

71 If the transducer has been beaten or knocked, please check whether the cover is airproof or damaged. If you have any doubt, please contact the manufacturer or local agent.

72 If the battery in the base station is stored alone and not used for a long time, we recommend that the battery should be charged at least once every 6 months to prevent over discharge.

73 The wireless transducer has priority over the wired transducer. When the wireless transducer is working, the wired transducer will be turned off automatically. Do not use the wireless transducer and the wired transducer at the same time.

CAUTION

- 1 The device is designed for continuous operation. Avoid liquid splashing on the device.
 - 2 Refer servicing to qualified personnel.
 - 3 Keep the environment clean. Avoid vibration. Keep it far from corrosive medicine, dust area, high-temperature and humid environment.
 - 4 When installing the unit into a cabinet, allow for adequate ventilation, accessibility for servicing, and room for adequate visualization and operation.
 - 5 Do not operate the unit if it is damp or wet because of condensation or spills. Avoid using the equipment immediately after moving it from a cold environment to a warm, humid location.
 - 6 The appliance coupler or mains plug is used as isolation means from supply mains. Position the monitor in a location where the operator can easily access the disconnection device.
-

CAUTION

- 7 Do not sterilize the monitor or any accessory with autoclave or gas.
- 8 Switch off the system power before cleaning. Cleaning consists of removing all dust from the exterior surface of the equipment with a soft brush or cloth.
- 9 Only the sensor and cable of US/TOCO transducers are watertight. Pay attention not let any liquid enter the transducer plug.
- 10 **Electromagnetic Interference** - Ensure that the environment in which the monitor or FTS-3 is installed is not subject to any source of strong electromagnetic interference, such as CT, radio transmitters, mobile phone base stations, etc. Even though other devices are in accordance with national standard radiation requirements, the monitor or FTS-3 may be interfered.
- 11 **Electromagnetic Interference** - Do not use mobile phones nearby in the process of monitoring
- 12 Operation of the equipment below the minimum amplitude may cause inaccurate results.
- 13 **Electromagnetic Interference** - Fetal parameters, especially ultrasound and ECG, are sensitive measurements involving small signals, and the monitoring equipment contains very sensitive high gain front-end amplifiers. Immunity levels for radiated RF electromagnetic fields and conducted disturbances induced by RF fields are subject to technological limitations. To ensure that external electromagnetic fields do not cause erroneous measurements, it is recommended to avoid the use of electrically radiating equipment in close proximity to these measurements.
- 14 **Electromagnetic Interference** - The monitor or FTS-3 system should not be used adjacent to or stacked with other equipment, refer to section A7.4 Recommended Separation Distances.
- 15 Electromagnetic interference is not unique to this system but is characteristic of fetal patient monitoring equipment in use today. This performance is due to very sensitive high gain front-end amplifiers required to process the small physiological signals from the patient. Among the various monitoring systems already in clinical use, interference from electromagnetic sources is rarely a problem.
- 16 The medical electrical equipment needs to be installed and put into service according to Appendix 7 EMC Information.
- 17 Portable and mobile RF communications equipment can affect medical electrical equipment, refer to section A7.4 Recommended Separation Distances.
- 18 Sterility cannot be guaranteed if package of the fetal spiral electrode is broken or opened.
- 19 The fetal spiral electrode has been sterilized by gamma radiation. Do not re-sterilize.
- 20 The device and reusable accessories could be sent back to the manufacturer for recycling or proper disposal after their useful lives.

CAUTION

21 If the terminals of the battery become dirty, wipe with a dry cloth before using the battery.

22 The device and accessories are to be disposed of according to local regulations after their useful lives. Alternatively, they can be returned to the dealer or the manufacturer for recycling or proper disposal. Batteries are hazardous waste. Do NOT dispose them together with house-hold garbage. At the end of their life hand the batteries over to the applicable collection points for the recycling of waste batteries. For more detailed information about recycling of this product or battery, please contact your local Civic Office, or the shop where you purchased the product.

In addition, when you use the FTS-3 fetal telemetry system, please pay attention to the cautions as follows:

1 The wireless transducers are IPX8 waterproof, but the base station should be kept non-soaked and non-condensing. The system may be condensing during transportation in high humidity or low temperature.

2 The water temperature must not exceed +60 °C (+140 °F) when you wash the belt.

3 The use of accessories and cables other than those specified may result in increased electromagnetic emissions or decreased electromagnetic immunity of the system.

4 This equipment generates, uses and radiates radio-frequency energy, and if it is not installed and used in accordance with its accompanying documentation, it may cause interference to radio communications.

5 When the battery is charged, used or stored, keep it away from objects or materials with static electric charges.

6 If the terminals of the battery become dirty, wipe with a dry cloth before using the battery.

7 The recommended charging temperature for the battery is between 0 °C ~ +40 °C (+32 °F ~ +104 °F). Please do not exceed the temperature range.

8 Batteries have life cycles. If the time that FTS-3 using battery becomes much shorter than usual, the battery life is at an end. Please contact the manufacturer to replace the battery with a new one of the same specification as the one provided or recommended by the manufacturer.

9 Remove the battery in the base station and store it at a cool and dry environment if the system is not used for a long time.

10 Please remove the battery out of the transducer at the end of its life.

11 Please read the user manual carefully when you install or remove the battery.

1.6 Definitions and Symbols

F9, F9 Express Fetal & Maternal Monitor

1		Socket for ultrasound transducer 1 (Type BF applied part)
2		Socket for ultrasound transducer 2 (Type BF applied part)
3		Socket for DECG cable (Type CF applied part)
4		Socket for TOCO transducer or IUP cable (Type BF applied part)
5		Socket for Remote Event Marker (Type BF applied part)
6		Socket for Fetal Stimulator (Type BF applied part)
7		Socket for NIBP Cuff (Type BF applied part)
8		Socket for SpO2 Transducer (Type BF applied part)
9		Socket for Maternal ECG Cable (Type CF applied part)
10		Socket for TEMP Transducer (Type CF applied part)
11		RS232 Interface (DB9 or D-Sub)
12		RJ45 Interface

13		Equipotentiality
14		Battery check
15		Alternating Current
16		Stand-by
17		Caution
18		Warning (Background: Yellow; Symbol & outline: Black)
19		Operating instructions
20		Refer to User Manual (Background: Blue; Symbol: White)
21		Type BF applied part
22		Defibrillation-proof type BF applied part
23		Type CF applied part
24		Defibrillation-proof type CF applied part
25	IPX1	Protected against vertically falling water drops
26	IPX8	Protected against the effects of continuous immersion in water
27		CE marking
28		Disposal method

29	P/N	Part Number
30		Date Of Manufacture
31		Manufacturer
32		Authorized Representative in the European Community
33		General symbol for recovery/recyclable
34	Rx Only	Caution: Federal (U.S.) Law restricts this device to sale by or on the order of a physician
35		This way up
36		Fragile
37		Keep away from rain
38		Stacking limit by number
39		Handle with care
40		Do not step on
FTS-3 Fetal Telemetry System		

35		Non-ionizing electromagnetic radiation
36		Serial Number
37		Wireless Transducer Working Indicator
38		USB Port (Reserved)
39		Ethernet Port (Reserved)
40		Channel Adjustment

NOTE:

The user manual is printed in black and white.

Chapter 2 Introducing the Monitor

2.1 Installation Guide

NOTE:

Installation must be carried out by qualified personnel authorized by the manufacturer.

2.1.1 Opening the Package and Checking

Visually examine the package prior to unpacking. If any signs of mishandling or damage are detected, contact the carrier to claim for damage.

Open the package; take out the monitor and accessories carefully. Keep the package for possible future transportation or storage. Check the components according to the packing list.

- ◆ Check for any mechanical damage.
- ◆ Check all the cables and accessories.

If there is any problem, contact us or your local distributor immediately.

2.1.2 Installing Battery

If your monitor has been configured with the rechargeable lithium-ion battery, follow these steps to install the battery:

WARNING

Switch off the monitor and unplug it before installing or removing the battery.

(1) Battery Installation

- a) Carefully place the monitor upside down on a flat surface covered with cloth or other type of protecting pad.
- b) Remove the screws of the battery compartment using a cross-head screw driver. Remove the battery compartment cover.



- c) Take the battery out from package. Put the battery and the cables into the battery compartment and insert the cable connector into the socket.



- d) Shut the battery compartment cover and fix the screws.



(2) Battery Removal

Fold the LCD display completely flat before turning the monitor upside down. Remove the battery in reverse order. To remove the battery, hold the two bands of the battery tight, shake it loose and pull it out with force.



NOTE:

- 1 If a rechargeable battery is outfitted, charge it fully each time after using the device to ensure the electric power is enough.
- 2 After the device is transported or stored for a long time, charge the battery fully before use. Connecting to power supply will charge the battery no matter if the monitor is powered on.
- 3 Do not pull the battery cables, or the battery may become damaged.

2.1.3 Installing Monitor

The monitor can be placed on a flat surface, or be installed on a wall or a trolley. The service engineer should install the monitor properly.

CAUTION

If you choose to install the monitor on the wall, MT-803 trolley, MT-503 trolley or other locations, it is the user's responsibility to ensure their integrity and solidity evaluated by a registered, professional structural or mechanical engineer and compliance with all local regulations. The manufacturer will not be responsible for the failure and loss of any improper installation.

2.1.4 Connecting Power Cable

- ◆ Make sure the AC power supply of the monitor complies with the following specification: 100V-240V~, 50Hz/60Hz.
- ◆ Apply the power cable provided with the monitor. Plug one end of the power cable to the power socket of the monitor. Connect the other end to a three-slot power output special for hospital usage.
- ◆ The equipotential grounding terminal is provided for the connection of a potential equalization conductor. Therefore, it is recommended to connect the grounding terminal of the monitor and the power outlet with the grounding wire, making sure the monitor is grounded.

WARNING

If the protective grounding (protective earth) system is doubtful, the power of the monitor must be supplied by internal power supply only.

NOTE:

- 1 Make sure the monitor and the power outlet are placed at a place where it is easy to connect and disconnect the power cord.
- 2 When the supply mains is interrupted, the device switches to internal power supply and operates normally if the battery is installed. If the battery is not installed, the

monitor shuts down and resumes the previous settings at the subsequent operation.

2.2 Overview

NOTE:

The pictures and interfaces in this manual are for reference only.



Figure 2-1 Appearance

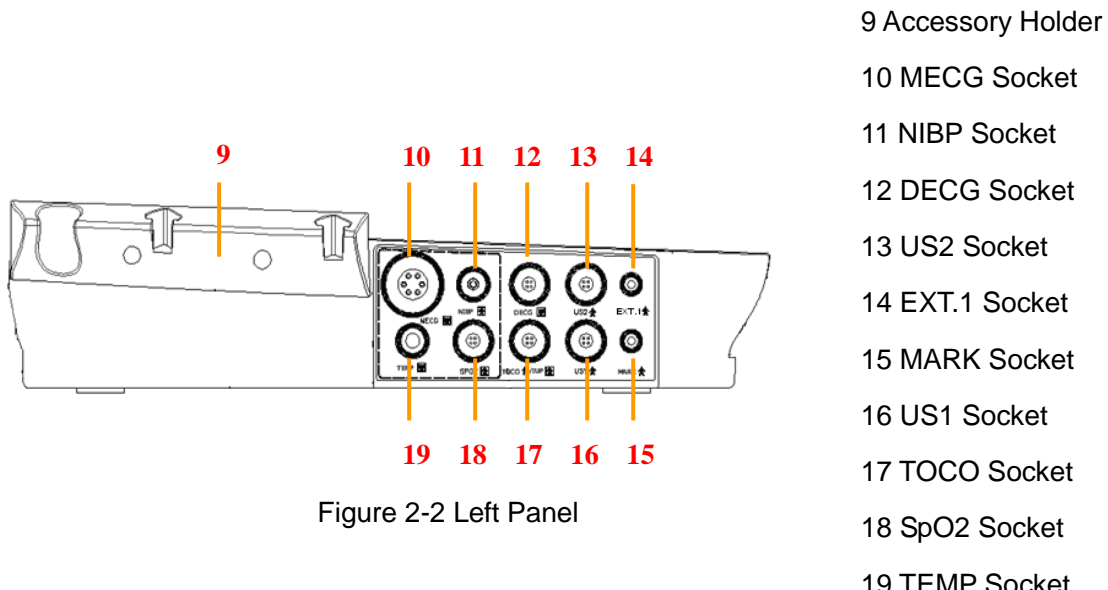


Figure 2-2 Left Panel

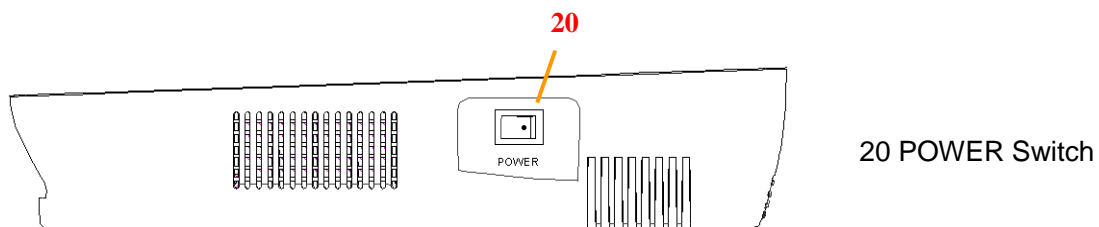


Figure 2-3 Right Panel

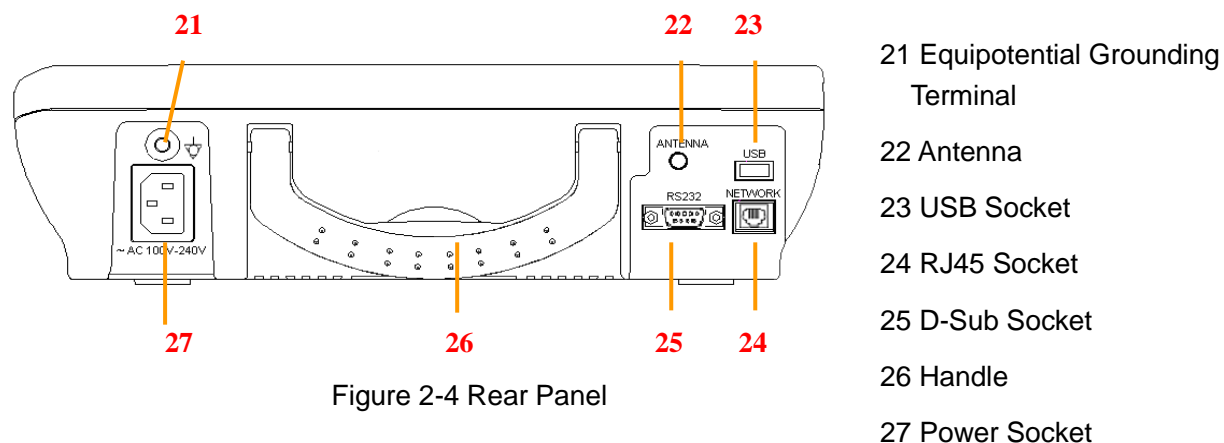


Figure 2-4 Rear Panel

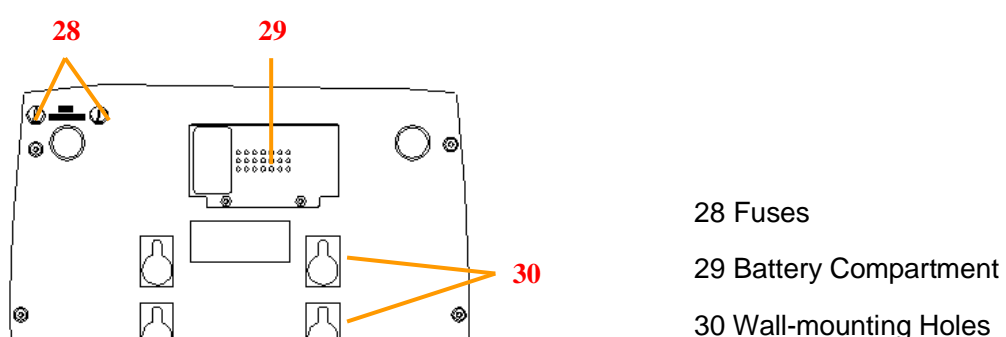


Figure 2-5 Bottom Panel

2.2.1 Keys

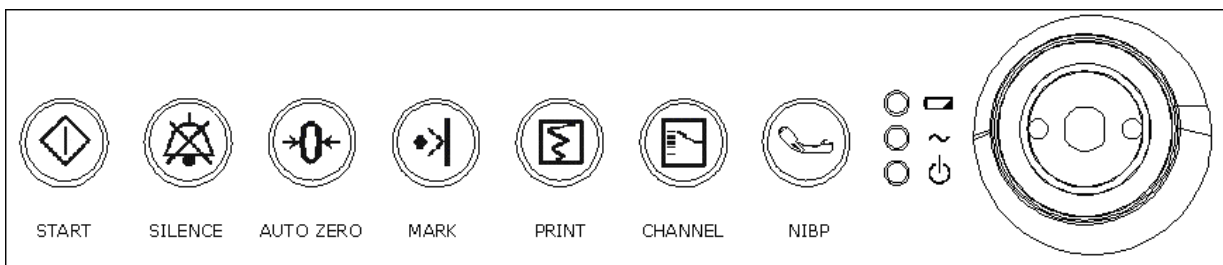


Figure 2-6 Keys and Control Knob

The Monitor is a user-friendly device with operation conducted by a few keys on the front panel and the control knob. Their functions are as follows:



Function: Start monitoring or return to the main interface

Press this key to start monitoring (on the main interface) or return to main interface (in information inputting menu, reviewing menu, searching menu or setup menus).



Function: Silence/reset

Press this key to disable the current auditory alarm manifestation, and re-enable the monitor's response to new abnormal patient condition.



Function: TOCO/IUP zero

Adjust the external TOCO contractions trace/numeric to preset unit (external monitoring contractions) or the IUP trace/numeric to reference point 0 (internal monitoring contractions).



Function: Record an event.

Press this key to make an event mark or open the smart note list.



Function: Start/stop printing

Press this key to toggle between starting and stopping printing.

(6) *CHANNEL



Function: Switch the channels

The monitor has two separate channels (channel 1 and channel 2). The default fetal heart sound comes from channel 1. When two transducers are connected to the monitor, press this key to switch the sound to channel 2; press it again to switch the sound back to channel 1.

(7) NIBP



Function: Start or stop a NIBP measurement.

Press this key to inflate the cuff and start a NIBP measurement. During the measuring process, press this key to cancel the measurement and deflate the cuff.

This function is only available on **F9 Express**.

(8) CONTROL KNOB

Function: Adjust volume, setup and playback control.

It can be rotated clockwise or counterclockwise and be pressed like other keys. All operations on the screen or in the menu can be completed by using the control knob.

The highlighted rectangular mark on the screen that moves with the rotation of the control knob is called “cursor”. Operations can be performed in the position on the screen where the cursor stays. When the cursor is located on a certain item, you can press the control knob to open its submenu or confirm the operation. Press the control knob again, and the cursor will be able to move around on the interface/menus.

Operating Procedure:

- Rotate the control knob to move the cursor to the required item.
- Press the control knob.
- One of the following three results will be achieved:
 - ◆ A new menu pops up. Operate the control knob in the new menu in the same way.
 - ◆ A submenu with several options appears on the right of the item. If this item has more than 8 options, they will be displayed in more than one page. Select **PREV** to switch to the previous page, or select **NEXT** to switch to the next page.
 - ◆ The function operates immediately.

**NOTE:**

- The word “select” hereinafter stands for rotating the control knob cursor to an item then pressing the knob.
- If the key sound is enabled, the monitor gives a normal key sound when the operation

is valid, and gives a sharp “Di” sound when the operation is invalid.

CAUTION

This monitor is a normal medical device. Please avoid violent operations such as continuously pressing the keys or control knob.

2.2.2 Indicators

There are four indicators on the top of the screen and the front panel. From the top down they are: alarm indicator, CHARGE indicator, AC indicator and Power indicator. The table below lists their meanings:

Indicator	Status of Indicator	Meaning
	Alarm Indicator	Flash or light up in yellow or red according to alarm level
		Off
 	Charge Indicator	On
		Off
 	AC Indicator	On
		Off
 	Power Indicator	On
		Off

2.3 Accessories

The accessories should be connected to the monitor via the sockets on the left side panel. Each accessory has a tab on the connector housing to ensure proper insertion into the appropriate socket on the monitor.

2.3.1 Ultrasound (US) Transducers



2.3.2 TOCO Transducers

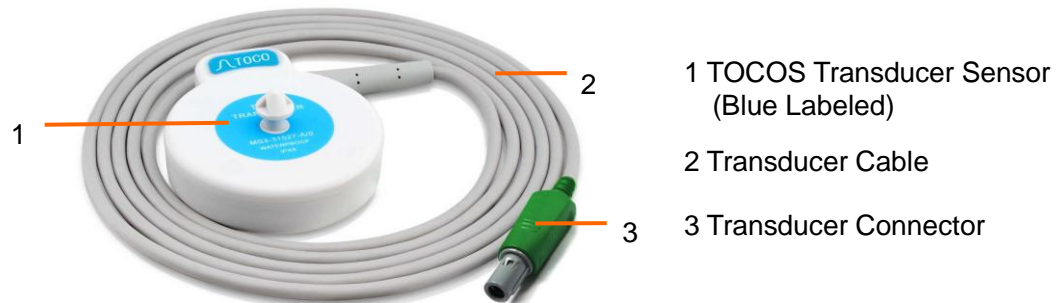


Figure 2-8 TOCO Transducers

CAUTION

Degree of protection against harmful ingress of water of the US transducer and TOCO transducer is IPX8. They can be continuously immersed in water to a depth of 1.1 meter for 24 hours and remain safe, but they are not allowed to be immersed in organic solvent, such as ethanol.

2.3.3 Belt



Figure 2-9 Belt

2.3.4 Remote Event Marker



Figure 2-10 Remote Event Marker

2.3.5 Fetal Stimulator

FS-1 Fetal Stimulator is a hand-held device. In order to reduce the time required for the NST when the fetus is asleep, it can be used to give a mild vibrating stimulation to the fetus through the maternal abdomen.

During NST, the vibrating operation marks can be displayed /printed on CTG trace when the fetal stimulator is connected to the monitor by an audio cable.

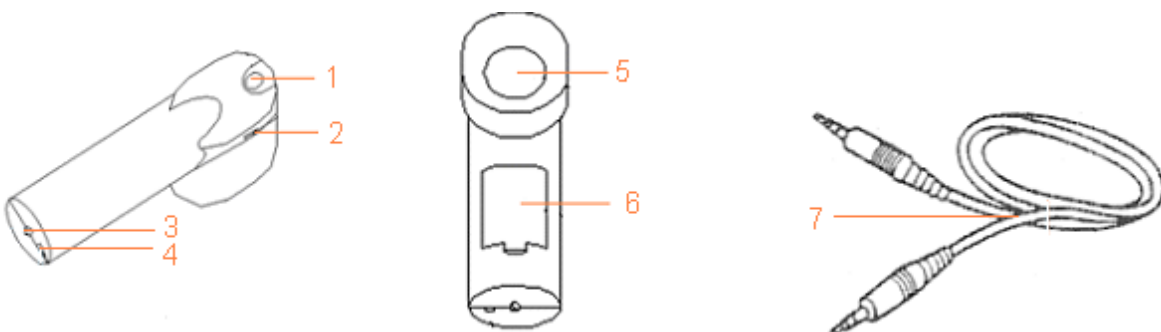


Figure 2-11 Fetal Stimulator

- | | |
|--------------------|------------------------------------|
| 1 Operating Switch | 2 Vibration Rhythm Adjusting Wheel |
| 3 Marker Socket | 4 Mode Selecting Switch |
| 5 Vibrating Head | 6 Battery Compartment |
| 7 Audio Cable | |

NOTE:

The fetal stimulator is NOT available in the USA.

2.3.6 DECG Cable



Figure 2-12 DECG Cable

2.3.7 Fetal Spiral Electrode

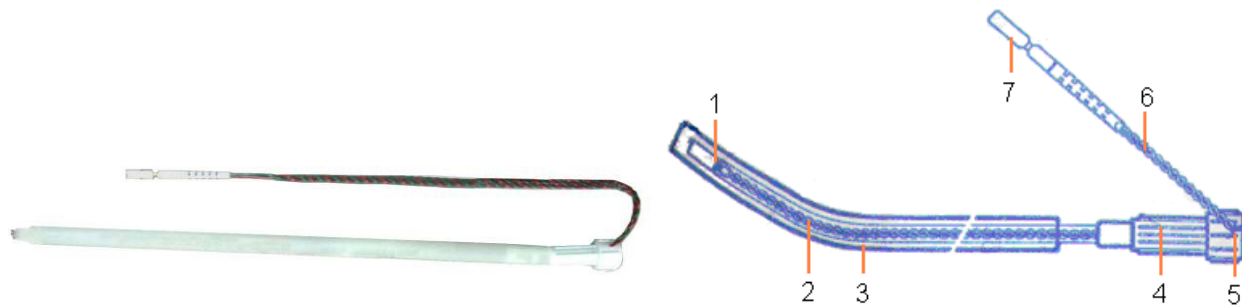


Figure 2-13 Fetal Spiral Electrode

- | | | | |
|-----------------------|------------------|--------------|----------------|
| 1 Reference Electrode | 2 Drive Tube | 3 Guide Tube | 4 Drive Handle |
| 5 Handle Notch | 6 Electrode Wire | 7 Safety Cap | |

2.3.8 IUP Cable



Figure 2-14 IUP Connecting Cable



Figure 2-15 IUP Cable

- | | |
|-----------------------------|---------------------------------|
| 1 Interface to IUP Cable | 2 Connecting plug |
| 3 Interface to IUP Catheter | 4 Interface to Connecting Cable |

2.3.9 IUP Catheter



Figure 2-16 IUP Catheter

2.3.10 ECG Cable



Figure 2-17 3-Lead ECG Cable

2.3.11 SpO₂ Sensor

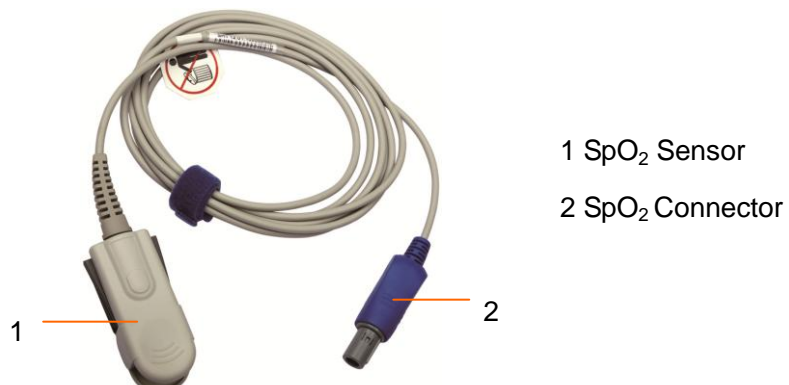


Figure 2-18 SpO₂ Transducer

2.3.12 NIBP Cuff



Figure 2-19 NIBP Cuff



Figure 2-20 Cuff Extension Tube

2.3.13 TEMP Sensor



Figure 2-21 TEMP Transducer

2.4 Screen

2.4.1 Main Interface

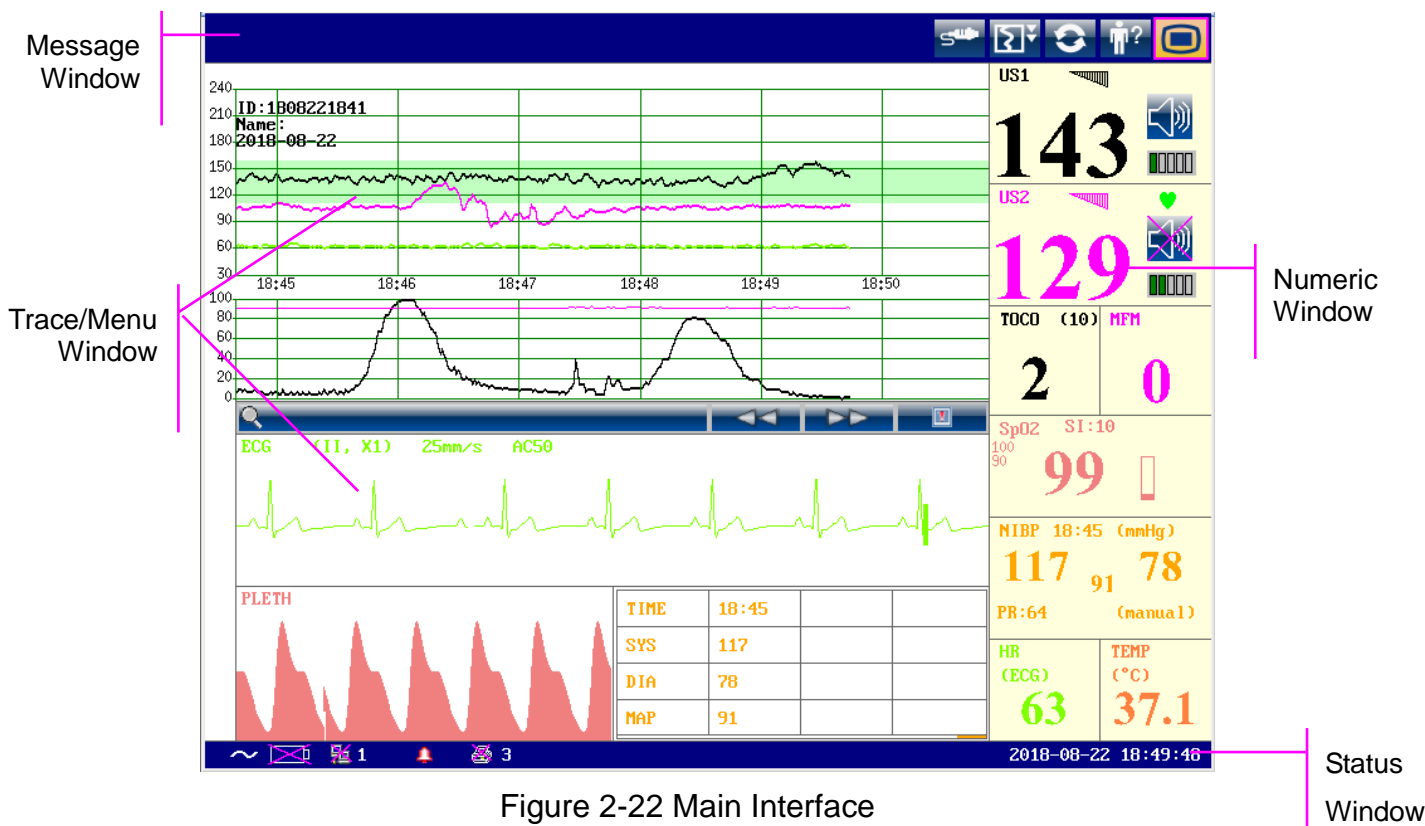


Figure 2-22 Main Interface

*Background Color Switch

The main interface of the monitor displays traces, waves, numerics, menus and monitor status information. The screen background color has four choices: black, green (default), orange and blue.

To change the screen color,

- 1 Select the setup key  on the main interface.
- 2 Select **General > Screen Color**.
- 3 Select the required color.
- 4 Select **OK**.

According to the content, the main interface is divided into four windows: (1) Message Window (2) Trace/ Menu Window (3) Numeric Window (4) Status Window.

(1) Message Window a



b c d e f





- a) **Alarm messages displaying area.** When an alarm is active, the message will be displayed here in yellow. Patient alarms will be displayed on the left and technical alarms in the center.



- b) **Confirming transducer unplugged alarm key.** When a transducer unplugged alarm and US1, US2 signal loss alarm is active, select this key to confirm the alarm and it will be turned off audibly and visually. But it still exists in the alarms review.

Note:

If **Confirming transducer unplugged alarm** key is turned off, then the icon will not be displayed.



- c) **Paper advancing key.** Select this key to advance the paper for 8 cm (PHILIPS paper) or 7 cm (GE paper).



- d) **Display mode switch.** **F9 Express** monitor has three display modes: maternal-fetal display mode, fetal display mode and maternal display mode. Select this key, and the display mode will switch to the next one in order.



- e) **Mat. Info key.** Select this key to open maternal information menu for inputting or changing the patient's ID and name.



- f) **Setup key.** Select this key to open setup main menu.

(2) Trace/Menu Window

The trace/menu window occupies most space of the screen. During monitoring or reviewing, it displays traces; during setting, it displays setup menus.

The background pane bar supports two standards: 30 ~ 240 (American standard) and 50 ~ 210 (International standard).

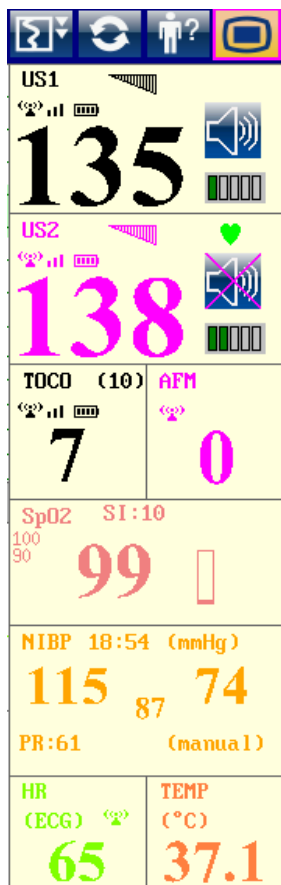
The green band in between the fetal heart rate panes indicates the preset alarm range (the top edge is not higher than 180 and the bottom edge is not lower than 100). It makes it easy to observe if the FHR exceeds the normal range. So you can easily tell if the fetal heart rate is too low or too high.



(3) Numeric Window

The fetal monitoring numerics and maternal vital signs are displayed here.

When the monitor is connected to the FTS-3 system, the signal strength icon , signal strength and quality icon , and battery level icon of the wireless transducers are displayed.



(4) Status Window



g) Power indicator

- AC power supplied.

- no AC power supplied.

h) Battery indicator

The battery is loaded into the monitor with 100% capacity

75% capacity

50% capacity

25% capacity

The battery is almost depleted and needs to recharge immediately.

No battery is loaded.

i) Network connection indicator and device no.

-  - the monitor is online.
-  - the monitor is offline.

NOTE:

The network connection indicator is not available if the net version is **Insight** or **Philips**.

- j) Audio alarm indicator
 -  - the audible alarm is switched on.
 -  - the current audible alarm is switched off infinitely.
 -  - the current audible alarm is switched off temporarily.
- k) Recorder status indicator
 -  - the recorder is in the process of printing.
 -  - no printing is going on.
- l)  - Print speed.
- m)  10min - Print remaining time.
- n)  ① 10:45 - Monitoring timer. It indicates the duration of the current monitoring, and zeroes when the **START** key is pressed.
- o)  « 12 - FTS-3 system working channel
- p) FTS-3 Base Station Battery indicator
 -  The battery is loaded into the base station with 100% capacity
 -  75% capacity
 -  50% capacity
 -  25% capacity
 -  The battery is almost depleted and needs to recharge immediately.

When there is no battery indicator, it indicates that no battery is installed in the base station.
- q)  2013-10-24 11:36:22 - Date and time of the monitor.

2.4.2 Setup Interface

Setup menus are provided to change the monitor configurations and monitoring settings. Press the setup key  on the main interface to open the main menu.

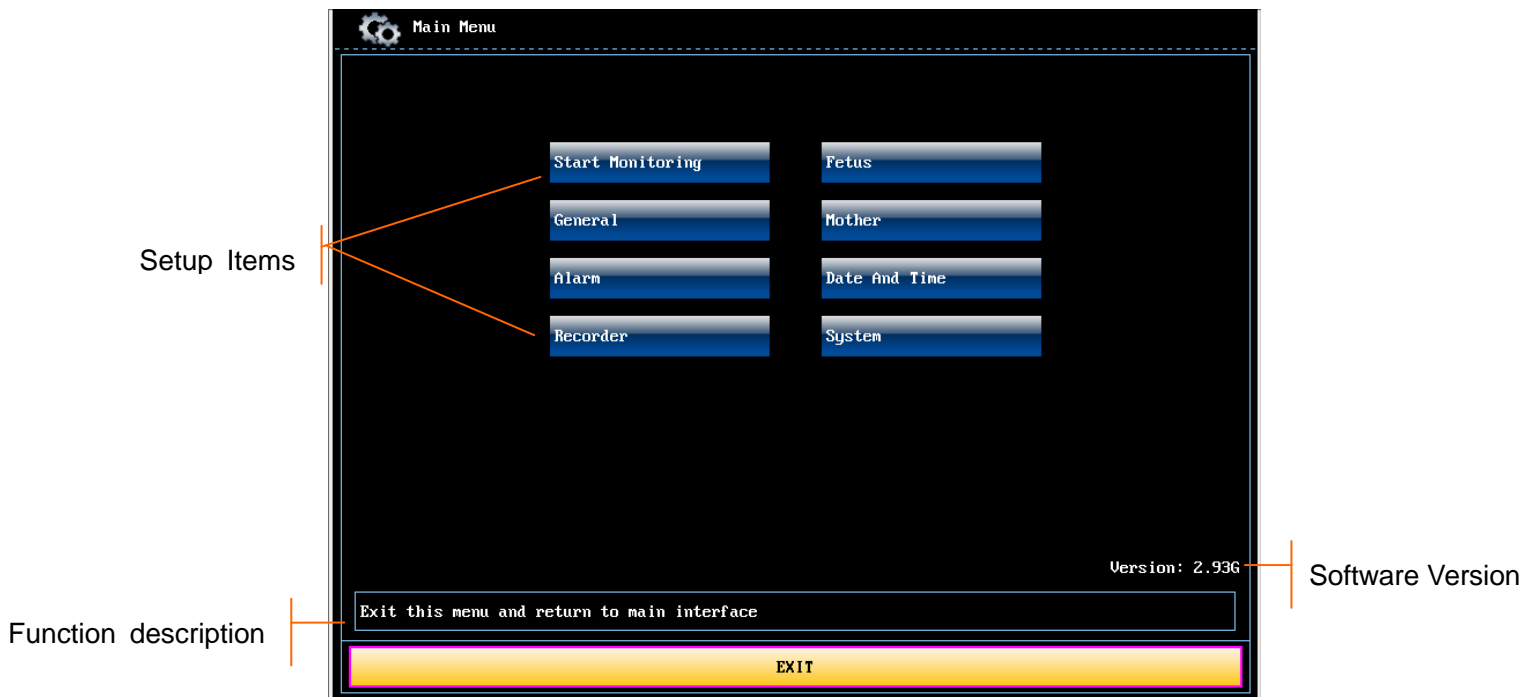


Figure 2-23 Setup Interface

In the setup main menu, you have access to all the items other than **System**. You can select **EXIT** to exit from this menu.

The items in this main menu all have submenu(s). To confirm the setting changes in the submenus, you need to select **OK** to exit. If you don't want to store the new settings, select **Cancel**, or press the **START** key to return to the main interface. If no operation is performed in 30 seconds, the menu will return to the upper directory. The change will not be stored.

Once you select **OK** to confirm the setting changes, the new settings will be stored in the monitor's long-term memory. If the monitor is switched on again after being switched off or a power loss, it will restore the new settings. The setting does not take effect if the system exits automatically or is shutdown before **OK** is selected.

For your reference, when the cursor is located at an item in this menu, the monitor provides a brief function description of this item in a pane with blue frame under the items. For example, the cursor is located at "System" in the illustration above. Correspondingly, its function "Set system items of the monitor" is issued in the blue frame pane.

2.4.3 Touch Screen

The touch screen is easy to use and operate. It works as a smart control knob. All the operations of the control knob can be done by gently touching the corresponding position on the screen.

When the touch screen is configured, touching the corresponding menu item is equal to rotating the control knob to this item and then pressing it. In the same way, one of the three results with the control knob will be achieved.

On the main interface, the symbols and might appear right next to the highlighted

item. Touch the  symbol to increase the numeric or move leftwards. While touching the  will decrease the numeric or move rightwards.

To exit from the submenu, you should touch the item again or touch any place outside the area of the options.

NOTE:

When touching an item, place the finger or the stylus pen within this item's cursor pane to ensure the operating validity. A key sound is heard corresponding to every valid touch, if the key sound is enabled.

Chapter 3 Introducing the FTS-3 Fetal Telemetry System

3.1 Brief Introduction

FTS-3 Fetal Telemetry System (hereinafter called FTS-3) provides monitoring for FHR, DECG MHR, AFM and TOCO for the pregnant women. When connected to a compatible fetal monitor, FTS-3 provides wireless patient monitoring in the antepartum period and during labor and delivery.

It is intended to be used only by trained and qualified personnel in antepartum examination rooms, labor and delivery rooms. It is not intended for use in intensive care units, operating rooms or for home use.

FTS-3 is used with F6, F9 series fetal/maternal monitor and connects to the monitor by signal cable. The wireless transducers monitor the FHR, TOCO parameters within certain distance, and then the base station sends them to the monitor through signal cable, and the monitor can display, alarm, print or review the parameters.

FTS-3 consists of the wireless US transducers (US-T transducers), the wireless TOCO transducer (TOCO-T transducer or TOCO-E transducer) and the base station.

The wireless signal can be transmitted in the Industrial Scientific Medical Band (ISM) according to the local regulations. The transmission range depends on where the system is used. It is recommended to use in hospital for better transmission. The transmission range is shorter in water than that in the air.

3.1.1 Base Station



Figure 3-1 Top Panel

	Name	Description
1	Docking Slot	Place, charge and manage the transducer.
2	Power Indicator	When you turn the power supply, the indicator is on.
3	AC Indicator	When AC power is supplied, the indicator is on.
4	Battery Indicator	When the base station battery is charging, the indicator is on. When the battery is in low level, it is flashing.
5	Wireless Connection Indicator	When the transducer connects to the base station successfully, the green light is on.
6	Charging Point	When you place the transducer in the docking slot, you can charge the transducer by these points.

WARNING

The charging point is specially used for charging the medical equipment and please do not touch the charging point and the patient at the same time.

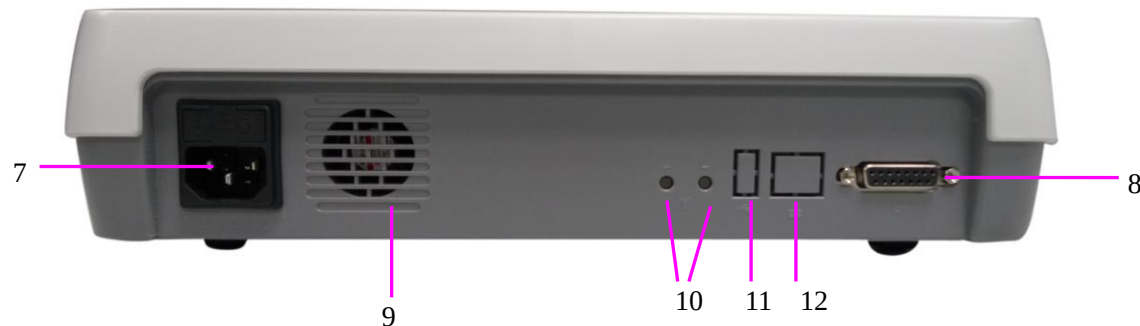


Figure 3-2 Rear Panel

	Name	Description
7	AC Outlet	AC outlet.
8	Communication Socket	Communicate to the bedside monitor.
9	Emission Slot	Emit the heat.
10	Channel Adjustment Button	Adjust the channel.
11	USB port	Reserved
12	Ethernet port	Reserved



Figure 3-3 Right Panel

	Name	Description
13	Power Switch	Turn on or turn off the base station.

CAUTION

1. This monitor is a normal medical device. Please avoid violent operations such as continuously pressing the power switch.
2. When the transducer is taken up, please do not power off the base station.



Figure 3-4 Bottom

	Name	Description
14	Battery Compartment	Install the battery.

3.1.2 Transducers



Figure 3-5 US-T transducer



Figure 3-6 TOCO-T transducer



Figure 3-7 TOCO-E transducer



	Name	Description
15	Transducer	Tied to the pregnant women.
16	Transducer Type	Indicate the transducer type.
17	System Working Channel	Indicate the system working channel.
18	Signal Indicator	Indicate wireless signal strength.

19	Battery Indicator	Indicate battery level.
20	TOCO-E Display	<ol style="list-style-type: none"> 1. Display “TOCO/DECG” when connected to DECG cable; 2. Display “TOCO/MECG” when connected to MECG cable; 3. Display “TOCO” when not connected to DECG cable or MECG cable.

3.1.3 Accessories

- FTS-3 DECG Cable

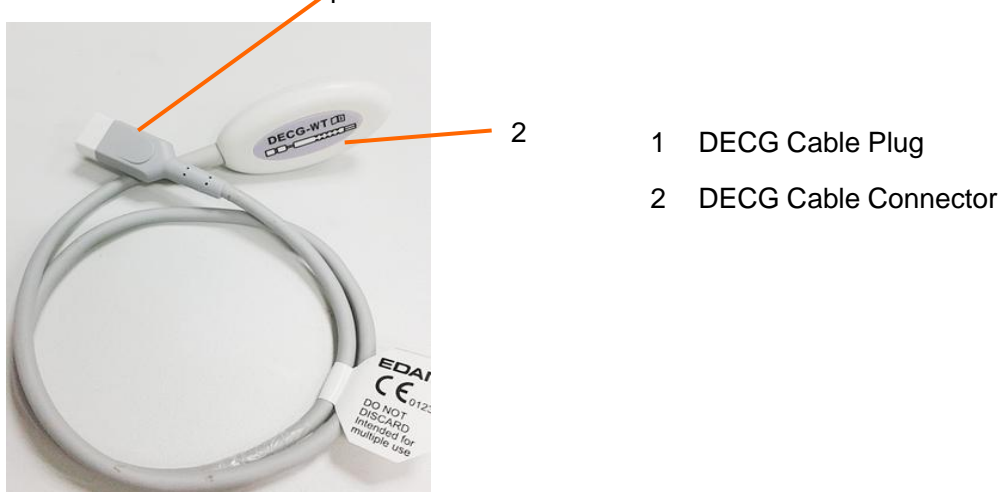


Figure 3-8 FTS-3 DECG Cable

WARNING

When connecting the fetal spiral electrode to the DECG cable, make sure that you have chosen the correct connector for the fetal spiral electrode according to the marking on the DECG cable.

- FTS-3 MECG Cable

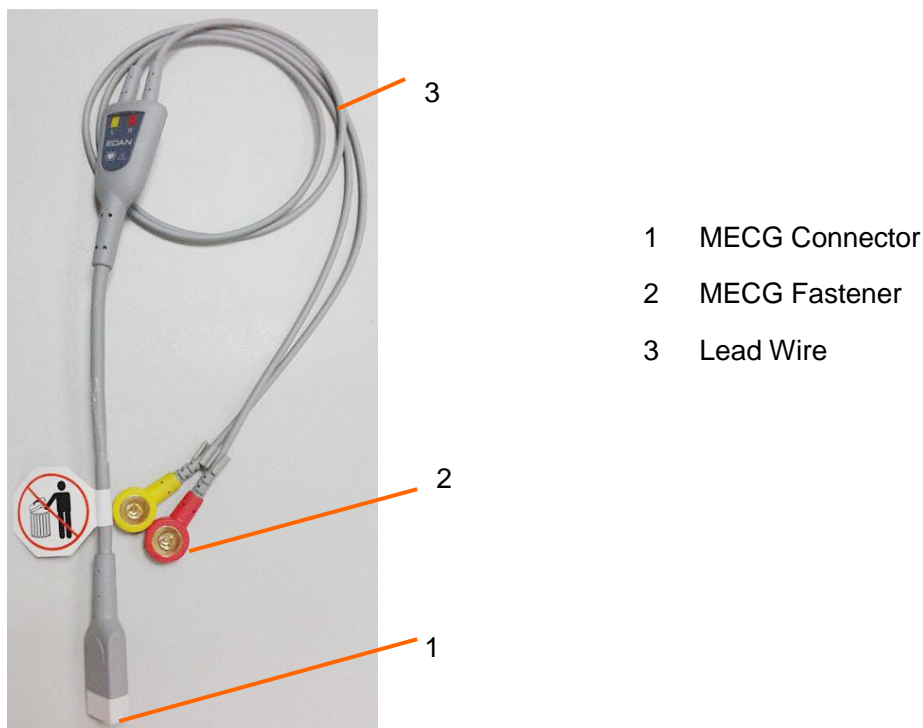


Figure 3-9 2-lead MECG Cable

3.1.4 Features

- Long work distance and free to walk in a great range
- Wireless transducers
- Low power consumption and working for a long time
- Rechargeable transducers
- Cabinet, portable and waterproof transducers
- Provide rechargeable battery for base station

3.2 Installation Guide

WARNING

The system installation should be operated by serviceman authorized by the manufacturer.

3.2.1 Opening the Package and Checking

Visually examine the package prior to unpacking. If any signs of mishandling or damage are detected, contact the carrier to claim for damage.

Open the package; take out the base station and accessories carefully. Keep the package for possible future transportation or storage. Check the components according to the packing list.

- ◆ Check for any mechanical damage.
- ◆ Check all the cables and accessories.

If there is any problem, contact us or your local distributor immediately.

3.2.2 Installing Battery

WARNING

Switch off FTS-3 and unplug it before installing or removing the battery.

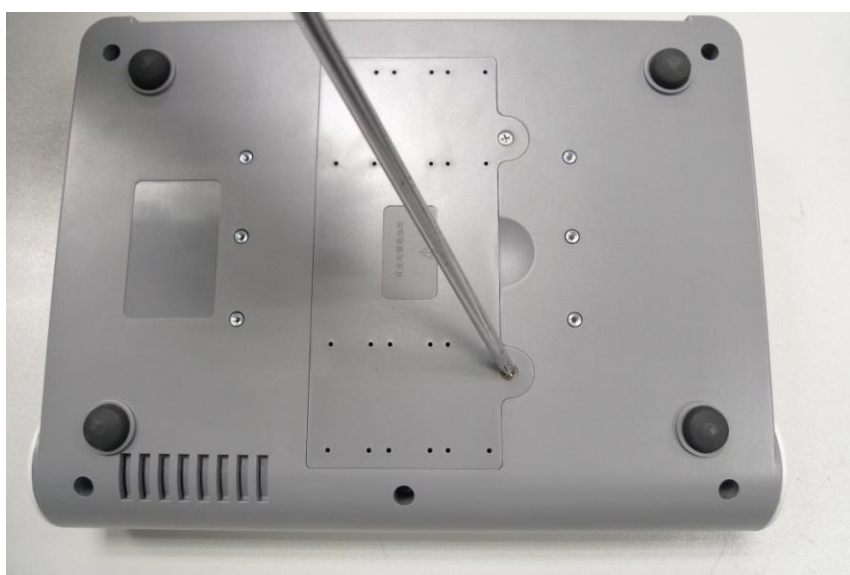
NOTE:

- 1 If the system is provided with a rechargeable base station battery, please charge the battery after each transportation and storage.
- 2 Please charge the battery to the full after each use. When the system is powered on with the AC power supply, the battery is charging. Please do not interrupt the charging and wait until the battery is fully charged.

If the system is provided with a rechargeable lithium-ion battery, follow these steps to install the battery:

(1) Battery Installation

- a) Place FTS-3 upside down on a flat surface covered with cloth or another type of protecting pad.
- b) Remove the screws of the battery compartment using a cross-head screw driver. Remove the battery compartment cover.



- c) Take the battery out from package and put it into the compartment. Make sure the battery connector is on the left and the battery label faces down.



WARNING

Do not touch the anode and cathode of the battery output together with fingers or metal materials, avoiding hazards to you and the battery caused by the short-circuit.

- d) Arrange the battery flat in the compartment, and push the strip at the end of the battery into the gap.



- e) Shut the battery compartment cover and fix it with the screws.

(2) Battery Removal

Remove the battery in reverse order. You can pull the strip at the end to take the battery out from the compartment.

NOTE:

- 1 If a rechargeable battery is outfitted, charge it fully each time after using the device to ensure the electric power is enough.
- 2 When the battery configuration is provided, after the device is transported or stored, the battery must be charged.

3.2.3 Installing the System

FTS-3 should be placed on a flat surface. It should be placed far from the device with strong radiation and avoid being in the shielded room. If there are 3 similar systems, they should be kept at a distance of over 1.5m from each other.

Alternatively, provided with proper devices, it can be installed on a wall or a trolley. Consult the sales representative for more information.

CAUTION

1. Installation must be carried out by qualified personnel authorized by the manufacturer.
 2. If you choose to install FTS-3 on the wall, MT-803 trolley, MT-503 trolley or other locations, it is the user's responsibility to ensure their integrity and solidity evaluated by a registered, professional structural or mechanical engineer and compliance with all local regulations. The manufacturer will not be responsible for the failure and loss of any improper installation.
-

3.2.4 Connecting Power Cable

- ◆ Make sure the AC power supply of the system complies with the following specification: 100V-240V~, 50Hz/60Hz.
- ◆ The equipotential grounding terminal is provided for the connection of a potential equalization conductor. Therefore, it is recommended to connect the grounding terminal of the system and the power outlet with the grounding wire, making sure FTS-3 is grounded.

WARNING

If the protective grounding (protective earth) system is doubtful, the power of the system must be supplied by internal power supply only.

NOTE:

- 1 Make sure the system and the power outlet are placed at a place where it is easy to connect and disconnect the power cord.

- 2 When the supply mains are interrupted, the device switches to internal power supply and operates normally if the battery is installed. If the battery is not installed, the system shuts down and resumes the previous settings at the subsequent operation.
- 3 After the AC power supply is connected, please wait for at least 2 seconds before pressing the POWER switch to turn on the system.

3.2.5 Configuration

The system can support, at most, 2 US-T transducers and 1 TOCO-T transducer, 2 US-T transducers and 1 TOCO-E transducer (DECG and MHR not enabled), 1 US-T transducer and 1 TOCO-E transducer connected with DECG cable, or 2 US-T transducers and 1 TOCO-E transducer connected with MECG cable. Please do not exceed the maximum number.

NOTE:

1. If the system is provided with transducer protection cover, please do not take up the cover during monitoring.
2. The TOCO-T transducer and the TOCO-E transducer cannot be used simultaneously.
3. When DECG is enabled, only US2 channel can be used for the US-T transducer. If one FHR is monitored by wired DECG and the other FHR is to be monitored by wireless US-T transducer, take up one US-T transducer, which will display "US1", and then wait for 2 seconds to take up the other US-T transducer. When the second US-T transducer connects successfully and displays "US2", fix it on the location where the best fetal heart signal is received, and put the first US-T transducer back in the docking slot.

3.2.6 Connecting to the Monitor

Use the DB15 interface cable (01.13.036299) to connect the FTS-3 system to the monitor. Then switch on the monitor and the FTS-3 system.

If the monitor is connected to the MFM-CNS Central Monitoring System, use the Y-shaped DB9 interface cable (01.13.036301) to connect the monitor and the FTS-3 system.

3.2.7 Adjusting the Working Channel

If the fetal heart sound is with interference or it cannot be played smoothly, the working channel is probably interfered. Put all the transducers back in the docking slots and press the adjustment

 button in back of the base station. The channel range is 1-14.

Restart the system when it enters the charging interface.

**NOTE:**

The working channel number used by a system cannot be duplicate with that used by a device of the same type.

3.3 Basic Operation

3.3.1 Charge the Transducer

Place the transducer in the docking slot and it displays the charging state on the transducer screen.

NOTE:

When the TOCO-E transducer is connected with the DECG or MECG cable, it cannot be charged due to poor contact with the charging points. Please remove the DECG or MECG cable before charging.

Caution

Please wait for 2 minutes to use the transducers after charging.

3.3.2 Charge the Battery

Please pay attention to the battery level during monitoring process. The battery symbol displays in the top right corner of the transducer screen. The low battery level may influence the monitoring.

: It is fully charged.

: It is less fully charged.

: It is in low level. Please charge the battery. There is alarm information on the screen.

: It is out of power. Please charge the battery immediately.

Caution

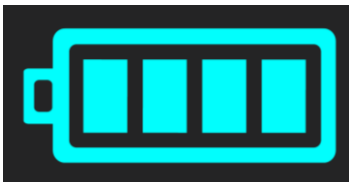
- 1 When indicates the power is low, please change for a new battery or charge the rechargeable battery in time, or the monitoring will be interrupted.
 - 2 After docking the transducer into the docking slot for charging, please check if the transducer is well placed and whether it is charging.
-

Please wipe the transducer and the charging point with a dry cloth before charging the transducer. Please do not scratch the charging point.

The battery is installed in the transducer. If the base station is switched on and supplied by AC, the battery will be charged automatically when it is placed in the docking slot. Please make sure the transducer is free of water and coupling gel before charging.

When you charge the battery, the screen will display as follows:

- Full charging icon: fully charged.



- Increasing charging icon: charging
- No charging icon: the transducer place in the docking slot incorrectly.
- If the screen displays ERROR, it indicates that the transducer is docked in the wrong base station.



The transducer has a screensaver function:

- When the transducer is fully charged in the docking slot for more than 10min ($\pm 1\text{min}$) and the base station is supplied by AC, a small yellow full charging icon will be displayed and floating on the screen and other icons will disappear.



- When the transducer is charging in the docking slot for more than 10min ($\pm 1\text{min}$) but the base station is not supplied by AC, a small yellow charging icon of real battery level will be displayed and floating on the screen and other icons will disappear.



- Take up the transducer or adjust the working channel to exit from the screensaver.

It takes about 3.5 hours to charge the battery. It is recommended to place the transducer in the docking slot when the transducer is not used for a long time.

Install the transducer in the base station and the transducer icon will display on the screen.

At the end of their life hand the batteries over to the applicable collection points for the recycling of waste batteries. If the battery charging time decreases sharply, the battery is considered as obsolete battery. Please use the battery provided by the manufacturer and disposes the battery according to the local regulations.

3.3.3 General Application

Take out the transducer from the docking slot and it will power on automatically. The transducer screen displays the signal strength, battery level and working channel. After the transducer is successfully connected to the base station, it will also display the transducer type. All the indicators are green. If the transducer is not successfully connected, it will power off automatically.

Take the transducer up and keep the transducer at a distance of over 30cm from the base station. The wireless connection indicator is on, and it indicates the transducer is taken out. If you want to power off the transducer, put it back in the docking slot. If the transducer connects to the base station successfully, the wireless connection indicator is always on and do not put back the inactivated transducer in the docking slot. Place the transducer on the patient.

WARNING

1. The DECG waveform, provided as a reference for signal quality, cannot be used for diagnosis. If the MECG cable is connected to the TOCO-E transducer and MECG monitoring is enabled, there will not be MECG waveform in the ECG display area of the main interface.
 2. The MECG cable is defibrillation-proof, but the DECG cable is not defibrillation-proof.
-

NOTE:

1. Detailed operations please refer to *7.2.2 Monitoring FHR with Wireless Ultrasound Transducer*, *7.3.5 Wireless DECG Monitoring*, *7.5.2 Wireless TOCO Monitoring* and *9.1.4 Wireless ECG Monitoring*.
2. If the working status indicator is on, please do not put the uncharged transducer in the docking slot.
3. Fix the US-T transducer, TOCO-T transducer and TOCO transducer tightly to ensure that they will not shift during movement.
4. It is recommended that the transducer should be placed when the patient stands for better monitoring.
5. Instruct the patient to move in the prescriptive area and distance for obtaining better signal.

6. When applied to the patient, the wireless ultrasound transducer may warm slightly (less than 3°C (5.4°F) above ambient temperature). When NOT applied, the wireless ultrasound transducer may warm slightly (less than 3°C (5.4°F) above ambient temperature).
7. The US-T transducer taken up first displays US1 on the screen, and the one taken up later displays US2. Please do not take two US transducers simultaneously and wait 2 seconds to take the other one. Restart the transducers if you take up two US-T transducers at the same time by mistake.
8. Please apply coupling gel to the US-T transducer before use and move the transducer to get the desired fetal heart and belt it to the belly. Underwater monitoring requires less coupling gel or no coupling gel. The TOCO-T transducer and TOCO-E transducer can be applied to the belly directly without coupling gel.
9. Excessive coupling gel may slide the transducer.
10. The TOCO-E transducer monitors DECG or MHR only when it is connected with the DECG or MECG cable. If the TOCO-E transducer is not connected with the DECG or MECG cable, it only monitors TOCO. Besides, the DECG cable and the MECG cable cannot be connected to the TOCO-E transducer at the same time.
11. When using the TOCO-E transducer to monitor DECG or MHR, it is recommended that the DECG cable or the MECG cable be kept straight to avoid damage to the TOCO-E transducer's interface caused by twisted cable.

3.3.4 Relocation of the Transducers

Transducers may be belted on the patient for a long time without stop. In rare cases, this may lead to irritations to the patient skin. To avoid skin irritations, please inspect the application site at least every half an hour. If the skin quality changes, you should move the transducer to another site.

US transducers need to change application site frequently to track fetal heart. It is normal during a monitoring process. But TOCO transducers are different. Please periodically inspect the application site (between contractions) of TOCO transducer at least every half an hour.

To reduce the risk of skin irritations, do not allow residual cleaning agent or disinfecting agent on the surface of transducers. Before using cleaning agent and/or disinfecting agent, refer to the cleaning and disinfecting sections in this user manual. Wipe the transducer surface with a cloth dampened with water before applying to the patient.

3.3.5 Monitor the Ambulatory Patient

FTS-3 is suitable for ambulatory patients. You can take out the transducer from the docking slot and fix the transducer on the location where the best fetal heart signal is received.

Please pay attention to the following during the monitoring.

- Ensure the transducer is tied up well.
- Record the effective FHR.

- The patient should not walk in strong tramps.
- The patient should move in the prescriptive area.
- The patient should be under monitoring when the wireless signal is good.

When the transducer is placed in the docking slot, the system stops transmission. It starts when the monitor is connected to the transducer.

When the patient moves during monitoring, the interference may occur. The artificial interference may influence the signal transmission quality. It will cause drop out or other interference if the transducer works in the changing environment. Some kind of the artificial interference can be anticipated and others can be discovered by observing the signal.

Some artificial interference may be caused by certain place. You can leave the place such as the elevator or the window in iron for the place with signal reception.

The FHR may not be detected clearly when the patient moves in virtue of artificial interference. The transducer is easy to shift underwater and it may lead to temporary signal loss.

No matter how good a telemetry system design is, the occasional US-T/TOCO-T/ TOCO-E dropouts are inevitable. If it is not acceptable for certain patients, please connect the wired the transducer to the bedside monitor.

The manufacturer has no control over the RF environment in the places where the system is used. If interference exists at operating frequencies, the system performance will be affected. You can change the working channel or move the system away from the interference to solve the problem.

Caution

- 1 The patient's steps may interfere with the monitoring of fetal heartbeats. It is suggested that the patient walks as less as possible.
 - 2 Excessive motion or vigorous movement may interfere with the monitoring and computing of FHR. Please try to avoid them.
-

3.3.6 Underwater Monitoring

Most wireless signal can be absorbed by water. Wireless transmission distances are shorter when monitoring under water. If you have any question, just contact the manufacturer or the local agent.

Caution

1. Please avoid flushing the transducer during underwater monitoring, or it may cause wireless signal interference.
 2. The transducers are watertight to a depth of 1.1 meter for 24 hours, but base station is not waterproof. Please do not splash water about the station or soak it into any liquid.
-

Caution

3. Underwater monitoring may influence the TOCO baseline in virtue of water temperature and depth or other reasons. Please adjust the TOCO baseline until the pressure of the transducer in water is steady and keep checking it.
 4. A metal bath tub and underwater monitoring both reduce the operating range.
 5. DECG and MHR cannot be monitored underwater.
 6. If the patient is monitored underwater, please place the transducers when she is ready.
-

3.3.7 Priority

- Priority in FHR1 numeric area: TOCO-E transducer with DECG cable > wired DECG cable > wireless US-T1 transducer > wired US1 transducer;
- Priority in FHR2 numeric area: wireless US-T2 transducer > wired US2 transducer;
- Priority in TOCO numeric area: wireless TOCO/TOCO-E transducer > wired TOCO transducer/IUP cable;
- Priority in MHR numeric area: TOCO-E transducer with MECG cable > wired MECG cable.

3.3.8 Basic Function Test

Please test the system after each service.

1. Power on the base station and connect it to the fetal monitor.
2. Charge the transducer.
3. Power on the monitor.
4. Take up the US-T transducer and test the following function:
 - The US-T transducer screen displays the standard start interface.
 - The US-T transducer indicator is green.
 - The fetal monitor screen displays US1 or US2.
5. Simulate the audio frequency signal:
 - The fetal monitor displays FHR.
6. Take up the TOCO-T or TOCO-E transducer and test the following function:
 - The TOCO-T or TOCO-E transducer screen displays the standard start interface.
 - The TOCO-T or TOCO-E transducer indicator is green.
 - The monitor screen displays TOCO.
7. Touch the measuring area of the TOCO-T or TOCO-E transducer gently:
 - The fetal monitor displays TOCO value change.
8. Install the US-T transducer to charge:
 - The US-T transducer screen displays charging interface and charging state.
 - The US-T transducer indicator is off.

- The fetal monitor screen has no display.
9. Install the TOCO-T or TOCO-E transducer to charge:
- The TOCO-T or TOCO-E transducer screen displays charging interface and charging state.
 - The TOCO-T or TOCO-E transducer indicator is off.
 - The fetal monitor screen has no display.
10. It takes about 3.5 hours to charge the US-T transducer, the TOCO-T transducer or the TOCO-E transducer.

Chapter 4 Alarms

WARNING

A potential hazard can exist if different alarm presets are used for the same or similar equipment in any single area.

4.1 Alarm Classification

The monitor has two types of alarm: patient alarm and technical alarm.

Patient alarms indicate the situation of vital sign exceeding its configured limit. Audible alarms and visual alarms can be disabled excluding ASYSTOLE alarm. The adjustable alarm limits determine the conditions that trigger the alarm.

Technical alarms indicate that the monitor cannot measure and therefore cannot detect critical patient conditions reliably. They cannot be disabled.

In terms of severity, the alarms are divided into three levels: high, medium and low. High level alarm indicates the condition where the patient's life is endangered; it is a severe warning, labeled with the symbol ***; Medium level alarm is a moderate warning, labeled with the symbol **; low level alarm is a general warning.

The high level alarms have highest priority, and the medium level alarms take the second place. If more than one type of alarms is active at the same time, the monitor sounds an audible indicator for the higher level alarms.

The alarm levels are preset, and you cannot change them.

4.2 Audible Alarm

If the audible alarm is not disabled, the alarm indicator displays . When an alarm is active, the monitor gives out a sound. (The sound pressure range is 45dB ~ 85dB.)

High level alarm: a "Do" tone is repeated three times, and then pauses for 3 seconds.

Medium level alarm: a "Do" tone is repeated three times, and then pauses for 5 seconds.

Low level alarm: a "Do" tone is issued, and then pauses for 20 seconds.

Press the **SILENCE** key, the current audible alarm toggles between on and off (temporarily or infinitely, you can change the setting).

If the current audible alarm is temporarily disabled, the alarm indicator displays SILENCE key is pressed.

If the current audible alarm is infinitely disabled, the alarm indicator displays 

is enabled again only when the **SILENCE** key is pressed.

If Alarm Reset is enabled (see *4.8 Pausing or Resetting the Alarm*), and you press the **SILENCE** key to disable an audible alarm, the alarm indicator will display , which is flashing in the manner of lighting for one second and pausing for one second. When other alarms present, the monitor will enable the audible alarm again automatically.

During the silence period, the alarm messages are displayed and the alarm indicator lights up as usual. You can press the **SILENCE** key again to enable the audible alarm.

WARNING

- 1 If the patient safety may be endangered, do not switch off the audible alarm infinitely.
- 2 Do not rely exclusively on the audible alarm system for patient monitoring. Adjustment of alarm volume to a low level or off during patient monitoring may result in patient danger. Remember that the most reliable method of patient monitoring combines close personal surveillance with correct operation of monitoring equipment.
- 3 When the sound pressure of audible alarm is equivalent to the ambient noise, it may be difficult for the operator to distinguish the audible alarm.

NOTE:

After you enable the audible alarm again, whether the alarm sound still exists depends on whether the alarm persists.

4.3 Visual Alarm

When an alarm is active,

- the alarm indicator lights up:

Alarm Category	Indicator Color	Flashing Frequency	Duty Cycle
High level alarm	red	1.4Hz to 2.8Hz	20% to 60% on
Medium level alarm	yellow	0.4Hz to 0.8Hz	20% to 60% on
Low level alarm	yellow	Constant (on)	100% on

- the alarm message appears in the message window of the main interface in yellow, with patient alarms on the left and technical alarms in the middle.

- the numeric of the measurement flashes in grey with a frequency of 2Hz.

When more than one alarm of the same level is active, the alarm messages appear in the same area in succession.

When more than one alarm of different levels are active, only the alarms of the highest level are displayed in the message window.

The patient alarm messages are displayed either:

- ◆ in text form, for example “** FHR2 LOW”; or
- ◆ in numeric form, for example “** FHR2 115 < 120”; ** indicates this is a medium level

alarm event; the first number is the current measurement result; the second number is the preset alarm limit.

The technical alarm messages are displayed in text form, for example “Fetus EQUIP MALF”.

WARNING

Setting alarm limits to extreme values may cause the alarm system to become ineffective. It is recommended to use the default settings.

4.4 Choosing the Alarm Display Form

You can change the patient alarm display form,

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Message Form**.
- 4 Select **Text** (default) or **Numeric**.
- 5 Select **OK**.

4.5 Changing the Alarm Volume

You can change the alarm volume,

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Alarm Volume**.
- 4 Select **Low** (default), **Medium** or **High**.
- 5 Select **OK**.

4.6 *Choosing Alarm Silence Duration

You can change the alarm silence duration,

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Silence Duration**.
- 4 Select **Infinite**, **1 min** (default), **2 min**, or **3 min**.
- 5 Select **OK**.

4.7 Choosing Signal Loss Delay

When the fetal signal is lost and this condition continues for a certain time, the monitor issues a technical alarm. This time (signal loss delay) is adjustable. To change the signal loss delay,

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Signal Loss Delay**.
- 4 Select **0** (default) ~ **300** seconds.
- 5 Select **OK**.

4.8 Pausing or Resetting the Alarm

You can enable the function of pausing or resetting audible alarms.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Audio Alarm**.
- 4 Select **Alarm Pause** (default) or **Alarm Reset**.
If **Alarm Pause** is selected: When the monitor gives out alarm sound and you press the **SILENCE** key, the alarm indicator displays , and the alarm sound is muted.
If **Alarm Reset** is selected: When the monitor gives out alarm sound and you press the **SILENCE** key, the alarm indicator displays , and the alarm sound is muted. When other alarms present, the monitor will enable the audible alarm again automatically.
- 5 Select **OK**.

4.9 Switching Transducer Unplugged On or Off

You can switch “Transducer Unplugged” on or off:

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, and then select **Enter**.
- 3 Select **Transducer unplugged** on the alarm settings interface.
- 4 Select **ON** or **OFF** (default).
- 5 Select **OK**.

4.10 *Reviewing Alarms

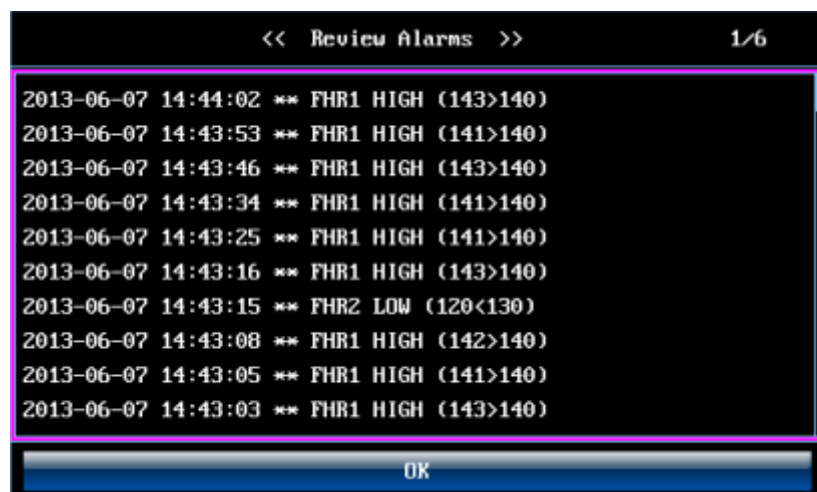
An alarm reviewing menu cannot only record the immediate alarm messages with date and time

information, but also record the historically physiological alarm, signal overlap alarm and US1, US2 signal loss alarm messages with date and time information.

The monitor can display a maximum of 100 immediate alarm messages. When the storage is full, it will delete the earliest alarm message automatically to store the new one.

The monitor can display a maximum of 800 historically physiological alarm, signal overlap alarm, and US1, US2 signal loss alarm messages. When the total number exceeds 800, the alarms messages cannot be stored.

Select the alarm reviewing key  to open this menu. When you review the traces with the word **REVIEW** shown in the background, the alarm reviewing menu displays historic alarm review. Otherwise, it displays the immediate alarm review.



Each page displays 10 alarm records. The page mark “1/6” informs you that there are 6 pages and the present one is page 1.

To review more records, select the alarm list and then rotate the control knob to switch to the previous or next page.

Select **OK** to exit from this menu.

When the monitor is switched off, the power supply is cut off accidentally, or a new monitoring starts, the immediate alarm messages will be cleared.

NOTE:

You can select **Main Menu > Alarm > Review Alarms** to set up **On** (by default) or **OFF**. When the alarm review is enabled, the icon will appear in the main interface.

4.11 Alarm Treatment Measures

During monitoring, make sure there is at least one physician in the area where the alarm sound can be heard or the alarm messages can be seen, so necessary measures can be taken when an

emergency occurs.

When the monitor gives out an alarm and catches your attention, you should:

- Check the patient's condition.
- Identify the cause of the alarm.
- Silence the alarm if necessary.
- Check if the alarm is terminated when the alarm condition is solved.

When the monitored parameter(s) come(s) back within the adjusted limits, or if the abnormal technical condition does not exist any longer, the monitor stops giving out the alarm.

If “Transducer Unplugged” is enabled (set to “ON”), and you press the confirming transducer

unplugged alarm key  on the main interface to confirm any active transducer unplugged alarms: US1 UNPLUGGED, US2 UNPLUGGED, TOCO UNPLUGGED, IUP UNPLUGGED, DECG UNPLUGGED, SpO2 SENSOR OFF, TEMP UNPLUGGED, US1 SIGNAL LOSS and US2 SIGNAL LOSS during the monitoring process, the transducer unplugged alarm(s) will be turned off audibly and visually until any of them occurs again. But the transducer unplugged alarm(s) still exist(s) in the alarms review list.

4.12 Testing Alarms

To test the functions of visible and audible alarms, do the following:

- 1 Switch on the monitor.
- 2 Enable the alarm.
- 3 Set the alarm limits to a small range.
- 4 Stimulate a signal that is higher than the upper limit or lower than the lower limit. Or disconnect one of the plugs.
- 5 Verify if the visible and audible alarms are working properly.

4.13 Patient Alarm Defaults

Alarm Setting	Options	Default
High Level		
ASYSTOLE	On (not adjustable)	On
Asystole Alarm Delay	0 seconds (not adjustable)	0 seconds
Asystole Alarm Level	High (not adjustable)	High
Medium Level		

FHR1/FHR2 Alarm	On, Off	On
FHR1/FHR2 Low Alarm Limit	60 bpm ~ 205 bpm, in increments of 5	110 bpm
FHR1/FHR2 High Alarm Limit	65 bpm ~ 210 bpm, in increments of 5	160 bpm
FHR1/FHR2 High Alarm Delay	0 ~ 300 second(s), in increments of 5	10 seconds
FHR1/FHR2 Low Alarm Delay	0 ~ 300 second(s), in increments of 5	10 seconds
FHR1/FHR2 Alarm Level	Medium, not adjustable	Medium
HR Alarm	On, Off	On
HR Low Alarm Limit	28 bpm ~ 242 bpm, in increments of 1	50 bpm
HR High Alarm Limit	29 bpm ~ 243 bpm, in increments of 1	120 bpm
HR Alarm Delay	0 second, not adjustable	0 second
HR Alarm Level	Medium, not adjustable	Medium
SpO ₂ Alarm	On, Off	On
SpO ₂ Low Alarm Limit	50% ~ 99%, in increments of 1	90%
SpO ₂ High Alarm Limit	51% ~ 100%, in increments of 1	100%
SpO ₂ Alarm Delay	0 second, not adjustable	0 second
SpO ₂ Alarm Level	Medium, not adjustable	Medium
SYS Alarm	On, Off	On
SYS Low Alarm Limit	40 mmHg ~ 269 mmHg, in increments of 1 (5.3 kPa ~ 35.9 kPa, in increments of 0.1)	90 mmHg (12.0 kPa)
SYS High Alarm Limit	41 mmHg ~ 270 mmHg, in increments of 1 (5.4 kPa ~ 36.0 kPa, in increments of 0.1)	160 mmHg (21.3 kPa)
SYS Alarm Delay	0 second, not adjustable	0 second
SYS Alarm Level	Medium, not adjustable	Medium
DIA Alarm	On, Off	On
DIA Low Alarm Limit	10 mmHg ~ 214 mmHg, in increments of 1 (1.3 kPa ~ 28.6 kPa, in increments of 0.1)	50 mmHg (6.8 kPa)
DIA High Alarm Limit	11 mmHg ~ 215 mmHg, in increments of 1 (1.4 kPa ~ 28.7 kPa, in increments of 0.1)	90 mmHg (12.0 kPa)
DIA Alarm Delay	0 second, not adjustable	0 second
DIA Alarm Level	Medium, not adjustable	Medium
MAP Alarm	On, Off	On
MAP Low Alarm Limit	20 mmHg ~ 234 mmHg, in increments of 1 (2.7 kPa ~ 31.2 kPa, in increments of 0.1)	60 mmHg (8.0 kPa)

MAP High Alarm Limit	21 mmHg ~ 235 mmHg, in increments of 1 (2.8 kPa ~31.3 kPa, in increments of 0.1)	110 mmHg (14.8 kPa)
MAP Alarm Delay	0 second, not adjustable	0 second
MAP Alarm Level	Medium, not adjustable	Medium
TEMP Alarm	On, Off	On
TEMP Low Alarm Limit	0 °C ~ +49.9 °C, in increments of 0.1	+36.0 °C
TEMP High Alarm Limit	+0.1 °C ~ +50.0 °C, in increments of 0.1	+39.0 °C
TEMP Alarm Delay	0 second, not adjustable	0 second
TEMP Alarm Level	Medium, not adjustable	Medium

NOTE:

The upper limit must be higher than the lower limit. When setting the upper limit, you do not have access to the options that are lower than the preset lower limit, and vice versa.

4.14 Alarm Messages

4.14.1 Fetal Monitoring Alarm Messages

During fetal monitoring, the monitor gives alarms for the situations that need the physicians to pay attention to. The alarm messages are listed below.

Patient Alarm Messages

Alarm Message	Source	Cause	Countermeasure
Medium Level			
**FHR1 HIGH or ** FHR1 xxx > yyy, **FHR2 HIGH or ** FHR2 xxx > yyy **DFHR HIGH or **DFHR xxx > yyy	US	FHR1, FHR2 or DECG measuring result (xxx) is higher than the set upper limit (yyy) and the alarm delay time of the upper limit is out.	Check if the alarm limits are suitable; check the patient's condition.
**FHR1 LOW or ** FHR1 xxx < yyy, **FHR2 LOW or ** FHR2 xxx < yyy **DFHR LOW or **DFHR xxx > yyy	US	FHR1, FHR2 or DECG measuring result (xxx) is lower than the set lower limit (yyy) and the alarm delay time of the lower limit is out.	

Technical Alarm Messages

Alarm Message	Source	Cause	Countermeasure

Medium Level			
**Battery Low	Monitor	The battery power is too low to support further work of the monitor.	Connect the monitor to AC power supply.
Low Level			
Check Paper	Monitor	There is no paper in the paper drawer or the drawer is open.	Load paper and/ or close the drawer.
US1 UNPLUGGED or US2 UNPLUGGED	US	US transducer 1 or US transducer 2 is not well connected. Or wireless US signal is not detected.	Check the connection of the transducer.
US1 SIGNAL LOSS or US2 SIGNAL LOSS	US	FHR1 or FHR2 signal is too weak for the system to analyze.	Check if the US transducer is aimed at the fetal heart; check if the alarm limits are suitable; check the patient's condition.
Fetus EQUIP MALF	US	The fetus board can not communicate with the system successfully.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
TOCO UNPLUGGED	TOCO	TOCO transducer is not well connected. Or wireless TOCO signal is not detected.	Check the connection of both TOCO transducer and US transducer.
IUP UNPLUGGED	IUP	IUP cable is not well connected to the monitor.	Check the connection of the IUP cable.
DECG LEADS OFF	DECG	The spiral electrode is not well connected.	Check the connection of the spiral electrode.
DECG UNPLUGGED	DECG	The DECG cable is not well connected to the monitor or the TOCO-E transducer.	Check the connection between the DECG cable and the monitor or the TOCO-E transducer.
DECG SIGNAL LOSS	DECG	DECG signal is too weak for the system to analyze.	Check if the spiral electrode is well attached to the fetus; check the patient's condition.
DECG EQUIP MALF	DECG	The DECG board cannot communicate with the system successfully.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
Signals Overlap (FHR1, FHR2)	US	US transducer 1 and US transducer 2 are aimed at the same fetal heart; the signals overlap.	Adjust one of the US transducers until another fetal heart signal is detected.

Signals Overlap (DFHR, FHR2)	US + DECG	US transducer 2 is aimed at the fetus that the spiral electrode is attached to; the signals overlap.	Adjust the US transducer until another fetal heart signal is detected.
Network Disconnection	Network	The central station disconnects with the fetal monitor (ETHERNET network)	Examine the network connection between the central station and the fetal monitor

4.14.2 Maternal Monitoring Alarm Messages

Besides the fetal monitoring alarms, **F9 Express** also gives alarms for the situations that occur during maternal monitoring. The alarm messages are listed below.

Patient Alarm Messages

Alarm Message	Source	Cause	Countermeasure
High Level			
***ASYSTOLE	ECG	No QRS wave is detected in 4 seconds	Check the patient's condition and take necessary measures.
Medium Level			
**HR HIGH or **HR xxx > yyy	ECG/ Pulse	Maternal HR result (xxx) is higher than the upper limit (yyy).	Check if the alarm limits are suitable; check the patient's condition.
**HR LOW or **HR xxx < yyy	ECG/ Pulse	Maternal HR result (xxx) is lower than the lower limit (yyy).	
** SpO ₂ HIGH or ** SpO ₂ xxx > yyy	SpO ₂	SpO ₂ result (xxx) is higher than the upper limit (yyy).	
** SpO ₂ LOW or ** SpO ₂ xxx < yyy	SpO ₂	SpO ₂ result (xxx) is lower than the lower limit (yyy).	
**SYS HIGH or **SYS xxx > yyy	NIBP	SYS result (xxx) is higher than the upper limit (yyy).	
**SYS LOW or **SYS xxx < yyy	NIBP	SYS result (xxx) is lower than the lower limit (yyy).	
**DIA HIGH or **DIA xxx > yyy	NIBP	DIA result (xxx) is higher than the upper limit (yyy).	
**DIA LOW or **DIA xxx < yyy	NIBP	DIA result (xxx) is lower than the lower limit (yyy).	
**MAP HIGH or **MAP xxx > yyy	NIBP	MAP result (xxx) is higher than the upper limit (yyy).	
**MAPLOW or **MAP xxx < yyy	NIBP	MAP result (xxx) is lower than the lower limit (yyy).	
**TEMP HIGH or **TEMP xxx > yyy	TEMP	TEMP result (xxx) is higher than the upper limit (yyy).	

**TEMP LOW or **TEMP xxx < yyy	TEMP	TEMP result (xxx) is lower than the lower limit (yyy).	
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Technical Alarm Messages

Alarm Message	Source	Cause	Countermeasure
High Level			
***ECG SIGNAL EXCEEDS LIMIT	ECG	ECG signal exceeds the measurement limits.	Check the connection of the leads and the patient's condition.
Low Level			
Signals Overlap (FHR1, HR)	US+ECG /Pulse	US transducer 1 has picked up the maternal heart signal; the signals overlap.	Reposition the US transducer 1 until the fetal heart signal is detected.
Signals Overlap (FHR2, HR)	US+ECG /Pulse	US transducer 2 has picked up the maternal heart signal; the signals overlap.	Reposition the US transducer 2 until the fetal heart signal is detected.
Signals Overlap (FHR1, FHR2, HR)	US+ECG /Pulse	US transducer 1 and US transducer 2 have picked up the maternal heart signal; the signals overlap.	Reposition the US transducers until the fetal heart signals are detected.
ECG LEADS OFF	ECG	The ECG cable is not well connected to the monitor or the TOCO-E transducer, or the leads of the ECG cable are not well attached to the patient.	Check the connection between the ECG cable and the monitor or the TOCO-E transducer. Check also the attachment of the ECG leads to the patient.
ECG EQUIP MALF	ECG	The ECG board cannot communicate with the system successfully.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
HR MEASUREMENT EXCEEDS RANGE	ECG/ Pulse	The heart rate exceeds the measurement limits.	Check the connection of the ECG leads/SpO ₂ transducer and the patient's condition.
NIBP EQUIP MALF	NIBP	The NIBP board cannot communicate with the system successfully.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
NIBP SYSTEM FAILURE	NIBP	The NIBP module defective.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
NIBP CUFF LOOSE or OFF	NIBP	The cuff is loose or not connected.	Wrap the cuff properly.

NIBP OVER PRESSURE	NIBP	The pressure has exceeded the specified upper safety limit.	Measure again, if failure persists, stop using the monitor for NIBP measuring and contact the manufacturer for service.
NIBP CUFF TYPE ERROR	NIBP	A different cuff other than the one supplied by the manufacturer is used.	Use the cuff supplied by the manufacturer.
NIBP LEAK	NIBP	The cuff, hose and (or) connector are (is) damaged.	Check and replace the leaking part(s). Contact the manufacturer for service if required.
NIBP SIGNAL LOSS	NIBP	Cuff is too loose or the patient pulse is too weak.	Use other methods to measure NIBP.
NIBP SIGNAL INTERFERED	NIBP	Large signal noise or irregular pulse rate caused by excessive motions of the patient.	Keep the arm that is wrapped with the cuff still.
NIBP EXCEEDS MEASUREMENT RANGE	NIBP	The blood pressure exceeds the measurement limits.	Check the connection of the cuff and the patient's condition.
NIBP TIME OUT	NIBP	Measuring time has exceeded 120 seconds.	Start measuring again, or use other measuring methods.
SpO ₂ LOW PERfusion	SpO ₂	The signal received by SpO ₂ sensor is too weak, or the measurement part has low perfusion, and therefore the result may be inaccurate.	Check the patient's condition and reposition the SpO ₂ sensor. Contact the manufacturer for service if the problem persists.
SpO ₂ SENSOR OFF	SpO ₂	SpO ₂ sensor is not well connected.	Check the connection of SpO ₂ sensor and finger placement.
SpO ₂ EQUIP MALF	SpO ₂	The SpO ₂ board can not communicate with the system successfully.	Restart the monitor and try again, contact the manufacturer if the connection still fails.
TEMP UNPLUGGED	TEMP	TEMP transducer is not well connected.	Check the connection of TEMP transducer.
TEMP EXCEEDS MEASUREMENT RANGE	TEMP	The temperature exceeds the measurement limits.	Check the connection of the TEMP transducer and the patient's condition.
TEMP Calibration Failed	TEMP	Calibration of the TEMP transducer failed.	Restart the monitor and try again. Contact the manufacturer for service if the problem persists.

4.14.3 FTS-3 Technical Alarm Messages

When FTS-3 is connected to F9 series fetal/maternal monitor, the monitor gives technical alarms for the situations that need the physicians to pay attention to during wireless monitoring. The alarm messages are listed below.

Alarm Message	Cause	Countermeasure
Medium Level		
**Wireless US1 Transducer Battery Low	The battery power is too low to support further work of the transducer.	Please charge the US1 transducer immediately.
** Wireless US2 Transducer Battery Low	The battery power is too low to support further work of the transducer.	Please charge the US2 transducer immediately.
** Wireless TOCO Transducer Battery Low	The battery power is too low to support further work of the wireless TOCO-T transducer or the wireless TOCO-E transducer.	Please charge the TOCO-T transducer or TOCO-E transducer immediately.
** Base Station Battery Low	The battery power is too low to support further work of the base station.	Connect the base station to AC power supply.
Low Level		
Wireless US1 SIGNAL LOSS	Wireless FHR1 signal is too weak.	Check if the patient moves out of the base station RF range, if the transducer is well connected to the base station.
Wireless US2 SIGNAL LOSS	Wireless FHR2 signal is too weak.	Check if the patient moves out of the base station RF range, if the transducer is well connected to the base station.
Wireless TOCO SIGNAL LOSS	The signal of the wireless TOCO-T transducer or wireless TOCO-E transducer is too weak.	Check if the patient moves out of the base station RF range, if the transducer is well connected to the base station.

4.14.4 Alarm Bilateral Control

When the network version is ETHERNET 1.4, alarm bilateral control is supported. The alarm settings that fetal monitor and central station synchronize bilaterally include high/low alarm limit, alarm switch and alarm delay of physiological alarm settings: FHR high/low alarm limit, FHR alarm switch and FHR alarm delay settings, HR high/low alarm limit, HR alarm

switch, SpO₂ high/low alarm limit, SpO₂ alarm switch, SYS high/low alarm limit, SYS alarm switch, DIA high/low alarm limit, DIA alarm switch, MAP high/low alarm limit, MAP alarm switch, TEMP high/low alarm limit, TEMP alarm switch.

When the fetal monitor gets on line, the central station will send above alarm settings to the fetal monitor for synchronization; the central station and the fetal monitor will send alarm settings to the other when alarm settings are revised on either of them after confirmation.

Chapter 5 Printing

5.1 *Function Description

The built-in thermal recorder applied in the monitor supports both the American and international standard wide recorder paper. It prints synchronously monitoring traces: FHR1/FHR2/TOCO/AFM/HR/ SpO₂ traces; trace marks: FHR1/FHR2/HR/ SpO₂; monitoring information: patient's ID, patient's name, date, time, speed (i.e. printing speed), FHR2 offset, HR, SpO₂, TEMP; SYS/DIA/MAP/PR; event marks: AUTO-zero symbol, MFM mark, event mark, fetal stimulation mark, alarm indicator, SOV indicator, US1 and US2 signal loss alarm indicator.

The monitor supports some other functions listed below:

- ◆ **Auto start printing:** If the function is enabled, the recorder starts printing automatically when new monitoring starts (the **START** key is pressed). Otherwise you have to press the **PRINT** key to start printing.
- ◆ **Printing timer:** The printing timer determines the elapsed time for each print. This time is adjustable. Refer to 5.2.3 *Changing the Print Timer*.
- ◆ **Remaining time indicating:** If the printing timer is set, a process indicator 10min appears in the status window after printing starts, with the remaining time shown in it. When the time is up, the monitor gives three “Do” tones and flashes the indicator.
- ◆ **Fast printing:** The recorder prints the data saved in the monitor at a high speed (up to 15mm/s).
- ◆ **Data Caching:** When the paper drawer runs out of paper or when it is open, the recorder stops printing. The data from this time on (at most 60 minutes) will be temporarily saved in the internal memory. When new paper is loaded and/or the drawer is closed, the saved data will be printed out at a high speed. When the saved trace has been printed out, the recorder switches back to continue printing the current data at the normal speed automatically.

NOTE:

- 1 When the monitor is switched off, the data in the internal memory will be lost.
 - 2 If a printing timer is set, and the time is out when the paper runs out, the CTG analysis result may disaccord with the printout. Therefore, reload the paper in time to avoid paper lack.
- ◆ **FHR2 offset:** You can set the offset of the FHR2 trace to separate the two FH traces on the screen and the recorder paper. Refer to 7.4.4 *Changing FHR2/DFHR Offset*.
 - ◆ **Print self-check:** The recorder prints a baseline for self checking when the monitor is switched on.
 - ◆ **Paper advance:** When printing stops, press the paper advancing key  to advance the paper, making sure the paper has a perforation outside the drawer and is easy to be torn off.

NOTE:

The paper advancing key is invalid in the process of printing and paper advancing.

5.2 Printing Configuration

NOTE:

All the parameters should be well configured before printing starts. You can not change the configuration in the process of printing.

5.2.1 Switching Auto Start Printing On or Off

You can switch auto start printing on or off:

- 1 Select the setup key  on the main interface.
- 2 Select **Start Monitoring > PRINT**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

5.2.2 *Choosing the Paper Speed

You can choose a paper speed of 1 cm/min, 2cm/min or 3cm/min:

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Print Speed**.
- 3 Select **1 cm/min, 2 cm/min** or **3 cm/min** (default).
- 4 Select **OK**.

NOTE:

Different paper speed setting causes different FHR trace appearance on the record paper. To avoid misinterpretation, we recommend you to set all monitors in your institution to the same paper speed.

5.2.3 *Changing the Print Timer

You can choose different time lengths for the print timer:

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Timer**.
- 3 Set timer to **10 ~ 90** (minutes, the step is 5) or **Infinite** (default). For a fixed time, the recorder stops when the time is up. For **Infinite**, there is no time limit. Whatever the setting is, the recorder stops when this patient's traces come to the end or if the **PRINT** key is pressed in midway.

- 4 Select **OK**.

5.2.4 Switching Print Self-Check On or Off

You can choose to switch print self-check on or off:

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Print Self-Check**.
- 3 Select **ON** (default) or **OFF**.
- 4 Select **OK**.

5.2.5 Changing Printing End Volume

The monitor gives a tone when printing ends and this tone volume is adjustable.

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Print Ending Beep**.
- 3 Select **High, Low** (default) or **OFF**.
- 4 Select **OK**.

5.2.6 Changing Title Print Cycle

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Title Print Cycle**.
- 3 Select **10min (Default), 20min, 30min, 60min**.
- 4 Select **OK**.

5.2.7 Switching Maternal Monitor Information On or Off

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > Maternal Monitor Information**.
- 3 Select **ON** (default) or **OFF**.
- 4 Select **OK**.

Note:

When the **Maternal Monitor Information** switch is turned off, the HR trace, SpO2 trace switch will be turned on automatically. You can turn them off manually if needed.

5.3 Understanding Recorder Paper Printout

WARNING

- 1 If there is any discrepancy between the display and the printout, the printout should prevail.
- 2 If the data is doubtful, clinicians should make diagnoses based on the real condition.

Figure 5-1 is an example of the recorder paper with traces. Comparing it with the monitor screen, you can find this extra information on it:

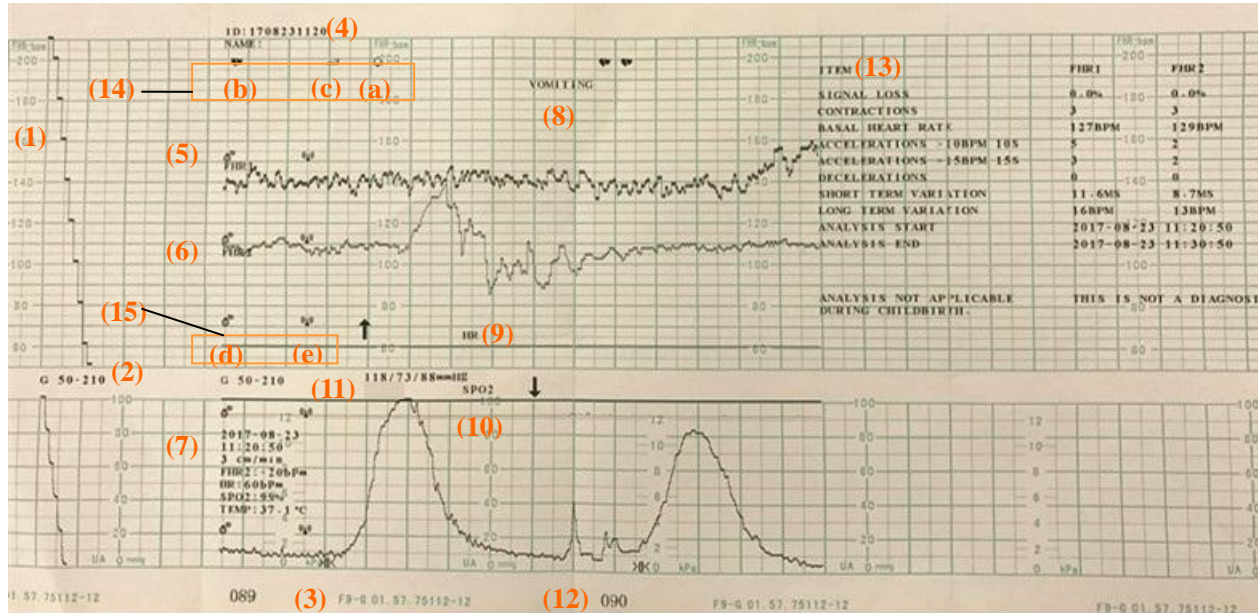


Figure 5-1 An example of recorder paper with traces

Item	Information	Description
1.	Self-Check Trace	The monitor prints a self-check trace after being switched on. It is used to check if the recorder paper is properly loaded.
2.	Paper Settings	The paper settings of the monitor. It consists of the paper type and paper style, e.g. "G 50-210", indicating that the paper type is "F9-G", and the paper style is International. It is printed out to check if the proper recorder paper is used.
3.	Paper Type	There are two types of paper: F9-G and F9-P.
4.	Paper Style	The FHR pane range indicates the paper style. American style: 30 ~ 240 International style: 50 ~ 210
5.	FHR1 Mark	The trace marked with "FHR1" is the FHR1 trace.
6.	FHR2 Mark	The trace marked with "FHR2" is the FHR2 trace.

7.	Trace Information List	A list of current date, time, print speed, ID, Name, FHR2 offset, HR, SpO ₂ and TEMP is printed at the start of the monitoring and every 10 minutes/20 minutes/30 minutes/ 60 minutes(optional) afterwards.
8.	Smart Note	The annotation of the event mark below.
9.	HR Mark	The trace marked with "HR" is the maternal HR trace.
10.	SpO ₂ Mark	The trace marked with "SpO ₂ " is the maternal SpO ₂ trace.
11.	NIBP	In the real-time printing mode, each NIBP measurement result is printed on the paper in the order of SYS/DIA/MAP/PR.
12.	Page Mark	Each recorder paper pack has 150 pages. When you notice the page mark comes to the end, remember to load new paper in time.
13.	CTG Analysis Result	The CTG analysis results of FHR1 and FHR2.
14.	Alarm Message	It indicates the physiological alarm message(a), signal overlap alarm message(b), US1 and US2 signal loss message(c).
15.	Wired and wireless monitoring mark	In the process of monitoring, if one channel changes between wired and wireless monitoring status, then wired mark (d) and wireless mark(e) will be printed according to the current data source of the channel.

Chapter 6 Pre-Monitoring Preparation

6.1 Loading Recorder paper

WARNING

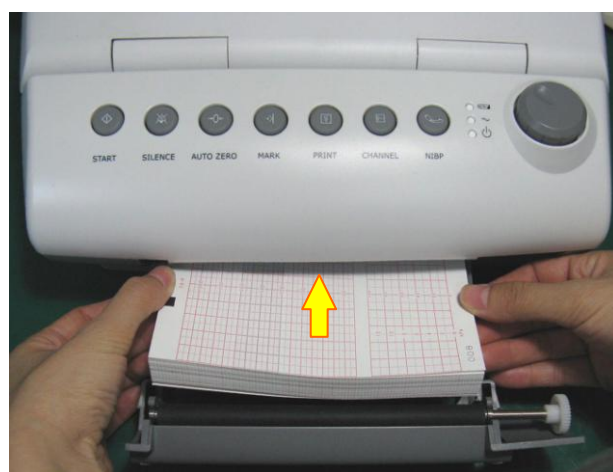
- 1 Only use the recorder paper provided by the manufacturer, otherwise the recorder may be damaged. This kind of damage is not covered by warranty.
- 2 Configured with different hardware, the monitor is compatible with both GE and Philips recorder paper. However, only one type of paper is configured with the monitor in the shipment. If you want to use the other type of paper, contact the manufacturer for service first, otherwise trace excursion or paper jam may occur.
- 3 Please check whether the recorder paper is correctly loaded.

If the monitor is used for the first time or when the paper runs out, you should load paper.

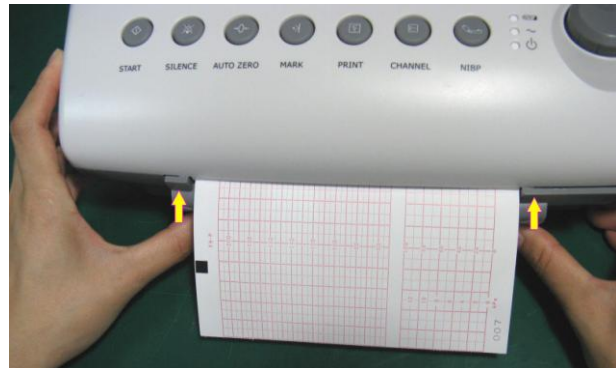
- 1) Press the two latches on each side of the paper drawer at the same time and slide the drawer out carefully.



- 2) Take out the Z-fold thermosensitive paper and remove the wrapper.
- 3) Place the pack in the drawer, with the pane facing up and the FHR trace area on the left.



- 4) Unfold two sheets from the top of the pack and pull the end of the paper out of the drawer (make sure the pack in the drawer remains flat).
- 5) Slide the drawer in until both the latches are locked.



NOTE:

- 1 Be careful when inserting paper. Avoid damaging the thermosensitive print head.
- 2 Make sure the paper is evenly loaded in the drawer. Otherwise the data will be inaccurate or paper jam will happen.
- 3 Only use the paper the manufacturer approved to avoid poor printing quality, deflection, or paper jam.
- 4 Keep the drawer closed unless when loading paper or servicing.

Removing Paper Jam

When the recorder does not function or sound properly, open the drawer to check for a paper jam. Remove the paper jam in this way:

- ◆ Cut the recorder paper from the paper drawer edge.
- ◆ Through the hole on the bottom panel of the paper drawer, push the recorder paper up with one finger. Remove the paper.
- ◆ Reload paper and then close the drawer.



6.2 Switching On the Monitor

WARNING

- 1 Check if all the metal parts are linked to the protective earth cord and the cord is in good condition before switching on the monitor.
- 2 If any sign of damage is detected, or the monitor displays some error messages, do not use it on any patient. Contact biomedical engineer in the hospital or our service engineer immediately.
- 3 Check all the functions to make sure that the monitor is in good condition.

Press the **POWER** switch on the right panel to switch on the monitor. The power indicator lights up and a start-up music will be heard. You can operate the monitor after the main interface appears.

You can choose to switch the start-up music on or off,

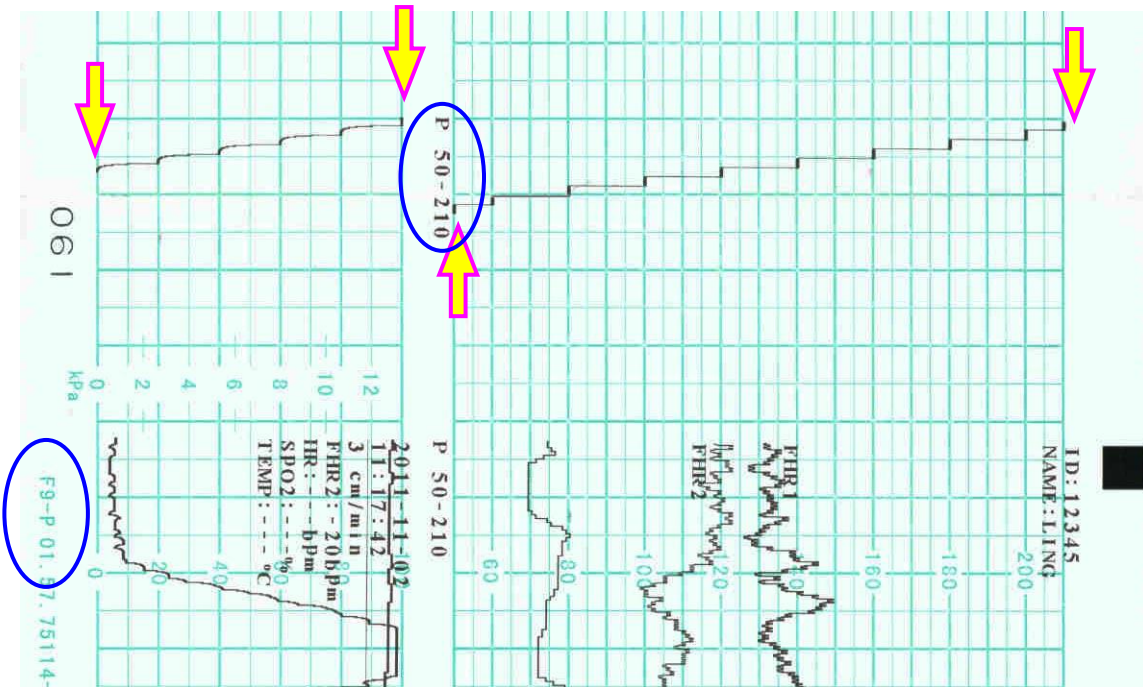
- 1 Select the setup key  on the main interface.
- 2 Select **General > Start-up Music**.
- 3 Select **ON** (default) or **OFF**.
- 4 Select **OK**.

6.3 Checking the Recorder Paper

The monitor provides the print self-check function to check if the recorder paper is correctly loaded and set.

The recorder prints a baseline and paper settings after start-up (if **Print Self-Check** is **ON**).

Check if the paper settings match the paper being used (in the circled area below, **P** should correspond to **F9-P**, and **G** to **F9-G**), and then observe the starts and ends of the printed baselines (illustrated with the arrow). The starts and ends should be printed exactly on the edges of the pane if the recorder paper is correctly loaded and set. If they do not comply with the edges, reload paper or ask the service engineer to check the paper settings of the monitor.



If the monitor does not print the baseline, switch on the **Print Self-Check** function and then restart the monitor.

NOTE:

Make sure the paper is correctly loaded before the printing starts.

6.4 Adjusting the Screen Angle

The angle between the screen and the top cover of the monitor is adjustable as needed, allowing it to be mounted on a wall or placed on a flat surface.

Adjustment method:

Push the hook on top of the screen left to spring it open. Pull the screen forward to let the screen stop at one of the three preset positions.



To bring the screen back to flat, pull it all the way forward and then push it back.



6.5 Setting Date and Time

You can change the date and time of the monitor,

- 1 Select the setup key  on the main interface.
- 2 Select **Date And Time**.
- 3 Set the year, month, date, hour, minute and second. The first three numbers are used to set the year, month and date. Their orders vary with the preset Date Format below.
- 4 Select **Date Format** for the format of the date; there are three options: yyyy-mm-dd (default), mm/dd/yyyy and dd/mm/yyyy.
- 5 Select **OK**.

CAUTION

- 1 You should set date and time information in advance. After this information is changed, the monitor starts new monitoring with an auto ID. Therefore, we advise you to restart the monitor after changing date or time information, and do not perform this operation when monitoring is in process.
- 2 If date and time cannot be saved, it is probably the battery has reached the end of its service life. Please contact the service personnel or your local distributor.

NOTE:

- 1 The date and time remain in the monitor for at least two months after it is switched off. You do not have to set date and time each time before monitoring.
- 2 When the network version is ETHERNET 1.4, if the fetal monitor gets on line or time is revised on the central station and the time difference between the fetal monitor and the central station is bigger than 1 minute, then the fetal monitor will synchronize time with the central station.

6.6 Connecting Transducers

Check for visible damages of the transducers every time before connecting them to the monitor. Pay special attention to the cracks on the transducers and cables before immersing them into conductive fluid. If damage is found, replace them with good ones at once.

When plugging transducers into the monitor, make sure the arrow symbol of the connector faces up and put it into the socket.



When disconnecting a transducer, pinch the afterbody of the transducer plug and pull it out slightly.

NOTE: Never try to disconnect the transducer by pulling the cable directly.

6.7 Placing Accessories in the Holder

In order to protect the accessories, place the not-in-use accessories in the holder. The accessory holder is on the left of the front panel. The first hole from the top is for the remote event marker, and the rest two are for the transducers.

To place a transducer into the holder, hold the transducer on the edge, and then place the buckle all the way into one of the holes on the holder. Make sure that the transducer cable is on the bottom.

To place the remote event marker, put the small end of the marker into the hole as far as it can go.

**NOTE:**

In the process of monitoring, the transducer that is placed in the holder may be affected and thereby produces interfering signals. Therefore, when monitoring a patient, it is recommended to remove or disconnect the transducer that is not in use.

6.8 Adjusting the Volume

The monitor automatically detects which channel the transducer is connected to. The corresponding volume adjustment key of this channel displays , indicating the FH sound is coming out from this channel; while the other one displays  , for example:



Press the **CHANNEL** key to switch the fetal heart sound to the other channel.

Adjust the default monitoring volume:

The FH volume returns to a default level after the **START** key is pressed. This default level is adjustable. To change this level,

- 1 Select the setup key  on the main interface.
- 2 Select **Start Monitoring > Volume**.
- 3 Select the volume from 1 ~ 10; the step is 1 and the default level is 3.
- 4 Select **OK**.

*Adjust the real-time monitoring volume:

If the default volume level is not satisfying during monitoring, you can adjust the real-time volume of each channel.

1 Select the volume adjustment key  on the main interface.

2 Rotate the control knob clockwise or touch the  symbol for one step, the volume increases by one level, there are ten levels in all; the green pane of the volume level indicator  increases by one at every two steps; rotate the knob anticlockwise or touch the  symbol to decrease the volume.

3 Press the knob again or touch any other place on the screen to confirm the volume level.

***Adjust the key volume:**

The volumes of pressing keys, rotating and pressing the control knob are also adjustable.

1 Select the setup key  on the main interface.

2 Select **General > Key Volume**.

3 Select **Low** (default), **High** or **OFF**.

4 Select **OK**.

Chapter 7 Fetal Monitoring

WARNING

- 1 The monitor is not intended for use in intensive care units (ICU), operating rooms or for home use.
- 2 Do not apply this monitor during electro-surgery or MRI; otherwise it might result in harming the patient or the operator.
- 3 Always check if the alarm settings are appropriate for your patient before starting monitoring.

7.1 Confirming Fetal Life

Fetal monitoring with ultrasound or DECG cannot differentiate a fetal heart rate signal source from a maternal heart rate source in all situations. These are some of the signal sources that might be taken as FHR signal source by mistake:

- High maternal heart rate signal.
- Maternal aorta or other large vessels signals.
- Electrical impulse from the maternal heart transmitted through a recently deceased fetus.

These are some of the signal sources that might be taken as fetal movement signal source by mistake:

- Movement of the ultrasound transducer.- Movement of the deceased fetus during or following manual palpation of fetal movement.
- Movement of the deceased fetus during or following maternal movement.

So you need to confirm fetal life by other means before starting to use the fetal monitor, such as using a fetoscope, stethoscope, Pinard stethoscope or obstetric ultrasonography.

7.2 Monitoring FHR with Ultrasound

The ultrasound monitoring is a method to obtain FHR on maternal abdominal wall. Place a US transducer (Ultrasound transducer) on maternal abdomen. It transmits low energy ultrasound wave to the fetal heart, and receives the echo signal. Monitoring FHR using ultrasound is recommended from the 24th week of gestation.

WARNING

- 1 Make sure you have confirmed the fetal life by other means before using this monitor for FHR monitoring.
- 2 FHR should not be monitored until a clear fetal heart signal is detected.

WARNING

- 3 If FHR reduces more than 10 bpm suddenly, or the beat of fetal heart sounds slower abruptly, please check if it is the MHR that is being monitored by the transducer. If so, relocate the transducer for the best fetal heart signal.
- 4 The sphere of activity for the fetus is much larger during mid-trimester of gestation (from 24th week to 28th week). When fetal heart moves away from the US transducer, please redetermine the fetal heart position and relocate the transducer.
- 5 Inspect patient skin before applying the transducer for monitoring. If the skin quality is poor, especially when the skin is damaged or irritated, please change a site to apply the transducer.
- 6 During long-time monitoring, please inspect the application site of the transducer at least every half an hour. If the skin quality changes, you should move the transducer to another site.
- 7 Do not mistake the maternal movement for fetal movement.

CAUTION

- 1 It is recommended to start printing FHR trace after clear fetal heart signal is detected and FHR computing has stabilized.
- 2 If AFM is enabled, suggest to reduce maternal movement and walking.

7.2.1 Monitoring FHR with Wired Ultrasound Transducer

7.2.1.1 Parts Required

- b) US transducer
- c) Aquasonic coupling gel
- d) Belt

7.2.1.2 Monitoring Procedure

a) Placing Transducer Belt

Place the transducer belts across the bed, ensuring that the belt will be around the abdomen when it is fastened. Lay the patient on the bed.

Alternatively, the patient can take a sitting position. Arrange the belt around her abdomen.

b) Determining the Transducer Position

- ◆ Determine the fetal position using Leopold's maneuvers.
- ◆ Search for the location of the fetal heart using a stethoscope or a fetoscope. The best fetal heart signal can be obtained through the fetal back.
- ◆ Place the transducer below the navel for head presentation and place the transducer

above the navel for breech presentation.

- ◆ During parturition, the fetal heart moves downward as the labor progresses. It is recommended to move the transducer along with the fetus.

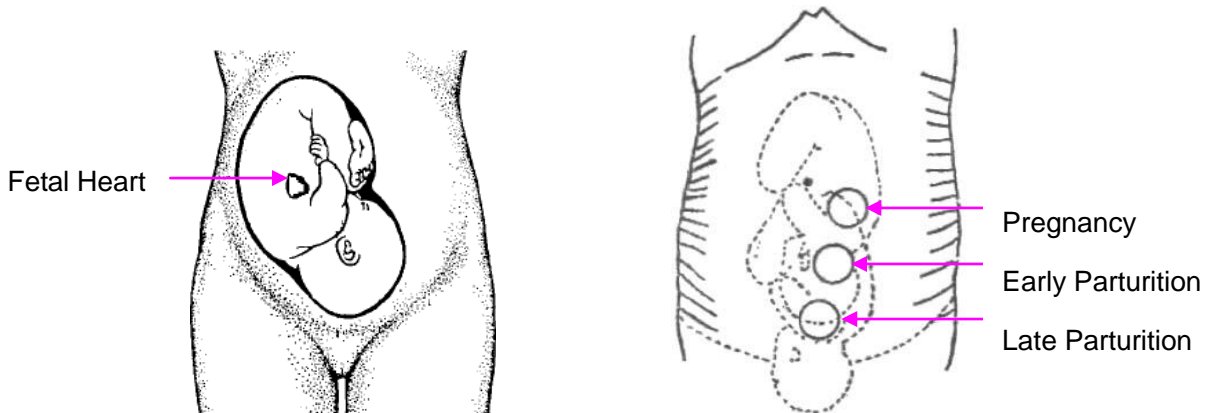


Figure 7-1 Positioning Ultrasound Transducer (single fetus)

c) Acquiring Fetal Heart Signal

Apply a certain amount of acoustic gel on the transducer and move the transducer slowly around the fetus site to even the gel. The best fetal heart signal can be obtained through the fetal back. Find at least 2 or 3 sites, and choose the one where the clearest, most sonorous and steady fetal heart sound is heard. When the transducer is connected correctly and communicated well, the fetal heart signal indicator is full. If the signal is poor, the signal indicator shows as it is and no FHR data are displayed. When you move the transducer on the abdomen, adjust the speaker volume so that it can be clearly heard.

d) Fixing the Transducer

When you find clearest and most steady fetal heart sound, wrap the abdomen with the belt over the transducer. Fix the transducer by pushing its buckle through the overlapping section of the belt.

Make sure the belt fits the patient snugly but comfortably. Meanwhile, fetus heart beat sound is heard; the FHR trace and numeric are displayed. During long-time monitoring, the gel may dry out as the transducer moves around. Add more gel in time if it is inadequate.

e) Confirming that the Fetus is the Signal Source

Ultrasound Doppler technology is utilized to observe the fetal heart rate externally when there are possibilities that maternal heart rate signal is mistaken for FHR signal. It is highly recommended to confirm that the fetus is the signal source continuously. You can perform either of the following:

- Measure the maternal heart rate with ECG or SpO₂ synchronously (available on **F9 Express** monitor). The monitor's SOV function can issue an alarm when the FHR signal source is likely to be from the maternal heart.
- Feel the maternal pulse at the same time.

If the maternal heart signal is misidentified as the fetal heart signal, Repositioning of the transducer is needed.

NOTE:

- 1 Do not mistake the high maternal heart rate for fetal heart rate. The fetal pulse can be distinguished from the maternal pulse by feeling the mother's pulse during the examination.
- 2 The best quality records will only be obtained if the probe is placed in the optimum position. Positions with strong placental sounds or umbilical blood flow sound should be avoided.
- 3 If the fetus is in the cephalic presentation and the mother is supine, the clearest heart sound will normally be found on the midline below the umbilicus. During monitoring, the patient's prolonged lying in the supine position should be avoided owing to the possibility of supine hypotension. Sitting up or lateral positions are preferable and may be more comfortable.
- 4 During long-time monitoring, the gel may dry out as the transducer moves around. Add more gel in time if it is inadequate.
- 5 When applied to the patient, the ultrasound transducer may warm slightly (less than 2°C (3.6°F) above ambient temperature). When NOT applied, at the ambient temperature of 40°C (104°F), the ultrasound transducer may reach the highest temperature of 43°C (109.4°F).
- 6 When an ultrasound transducer is connected to the monitor, but not applied to the patient, the measurement may generate unexpected intermittent FHR readings.

7.2.2 Monitoring FHR with Wireless Ultrasound Transducer

7.2.2.1 Parts Required

- a) US-T transducer
- b) Aquasonic coupling gel
- c) Belt

7.2.2.2 Monitoring Procedure

- a) Connect the FTS-3 system to the monitor and switch on. Refer to section 3.2.7 *connecting to the Monitor*.
- b) Take up a US-T transducer and make sure it is successfully connected to the base station.
- c) Monitor FHR following the procedures described in section 7.2.1.2 *Monitoring Procedure*.

7.2.3 Switching the FHR Alarm On or Off

Always check if the alarm settings are appropriate for your patient before starting a monitoring.

You can choose to switch the FHR alarm on or off. If the fetal heart alarm is switched off, the

monitor will no longer give any audible or visual warning for this monitoring item.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **FHR**.
- 4 Select **ON** (default) or **OFF** for **Alarm**.
- 5 Select **OK**.

If FHR alarm is switched off, an alarm switched-off symbol  will appear in the numeric window. For example:



WARNING

Do not switch the alarm off for the condition where the patient's safety maybe endangered.

7.2.4 Changing the FHR Alarm Limits

You can change the FHR alarm limits. The alarm limits you set determine the conditions that trigger the alarm.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **FHR**.
- 4 Select a value from 65 ~ 210 for **High Alarm Limit**. (The step is 5, and the default value is 160 bpm.)
- 5 Select a value from 60 ~ 205 for **Low Alarm Limit**. (The step is 5, and the default value is 110 bpm.)
- 6 Select **OK**.

7.2.5 Changing the FHR Alarm Delay

You can change the FHR alarm delay. The alarm delay indicates how long the measured result continues exceeding its limit before the alarm is triggered.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **FHR**.

- 4 Select a value from 0 ~ 300 second(s) for **High Alarm Delay**. (The step is 5, and the default value is 10 seconds.)
- 5 Select a value from 0 ~ 300 second(s) for **Low Alarm Delay**. (The step is 5, and the default value is 10 seconds.)
- 6 Select **OK**.

WARNING

The FHR alarm delay is adjustable between 0 and 300 seconds.

7.2.6 Testing US Transducers

To test a US transducer:

- 1 Switch on the monitor.
- 2 Connect the US transducer to the fetal monitor.
- 3 Hold the transducer with one hand, and gently touch the center of the transducer with the other hand in the frequency of 2 times per second.

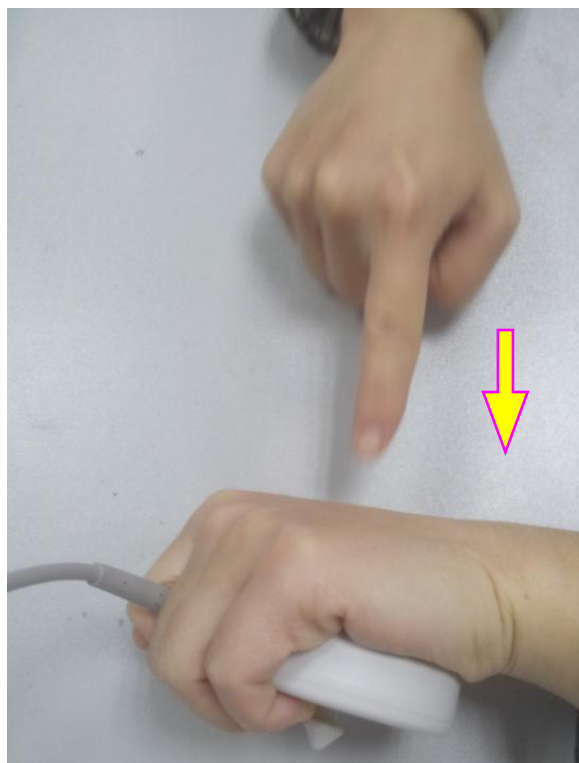


Figure 7-2 Testing a US Transducer

- 4 Check that the value on the display shows this change in FHR.

If a US transducer fails the test, repeat this test with another transducer. If the second one passes the test, defect of the first transducer is confirmed. Replace it with a good one. If the second transducer fails the test as well, contact the manufacturer for service.

7.3 Monitoring FHR with DECG

7.3.1 Contraindications

The fetal spiral electrode can be used when amniotic membranes are adequately ruptured and sufficient cervical dilatation is ensured. The fetal electrode tip is designed to penetrate the epidermis of the fetus; therefore, trauma, hemorrhage and/or infection can occur. The electrode should be used with strict adherence to aseptic technique.

The fetal spiral electrode should not be applied to the fetal face, fontanelles or genitalia.

Do not apply the fetal spiral electrode when placenta previa is present; when the mother has visible genital herpes lesions or reports symptoms of prodromal lesions; when the mother is HIV sero-positive; when mother is a confirmed carrier of hemophilia and the fetus is affected or of unknown status; or when it is not possible to identify fetal presenting part where application is being considered. This method is not recommended when fetus is extremely premature, or in the presence of a maternal infection such as Hepatitis B, Group B hemolytic strep, syphilis or gonorrhea, unless a clear benefit to the fetus or mother can be established.

7.3.2 Preparing the Patient's Skin Prior to Placing Electrodes

The skin is a poor conductor of electricity; therefore preparation of the patient's skin is important to facilitate good electrode contact to skin.

- ◆ Shave hair from electrode sites, if necessary.
- ◆ Wash the sites thoroughly with soap and water. (Do not use ether or pure alcohol, which will increase skin impedance)
- ◆ Rub the skin briskly to increase capillary blood flow in the tissues.
- ◆ Remove skin scurf and grease.

7.3.3 Directions for Using Fetal Spiral Electrode

- 1 With the patient in the dorsal lithotomy position, perform a vaginal examination and clearly identify the fetal presenting part.
- 2 Remove the spiral electrode from the package; leave the electrode wires locked in the handle notch.
- 3 Gently bend the guide tube to the desired angle.
- 4 Hold the drive handle, ensure the spiral electrode is retracted about one inch (2.5 cm) from the distal end of the guide tube.
- 5 Place the guide tube firmly against the identified presenting part.
- 6 Maintain pressure against the fetal presenting part with guide and drive tubes. Rotate the drive tube by rotating the drive handle clockwise until gentle resistance is encountered. Resistance to further rotation and recoil of the drive handle indicates that the spiral electrode

- is well attached to the fetus.
- 7 Release the electrode wires from the handle notch and straighten them. Slide the drive and guide tubes off the electrode wires.
 - 8 Insert the safety cap into DECG cable.

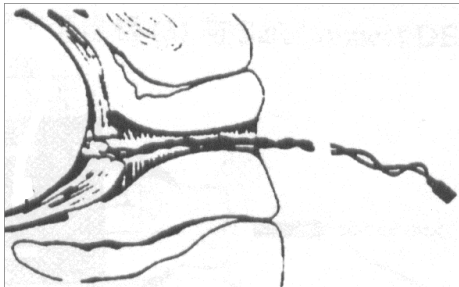


Figure 7-3 The Well-Attached Fetal Spiral Electrode

7.3.4 Wired DECG Monitoring

7.3.4.1 Parts Required

- a) DECG cable
- b) Fetal spiral electrode
- c) Disposable maternal attachment pad electrode

The following illustration shows how these parts should be connected:

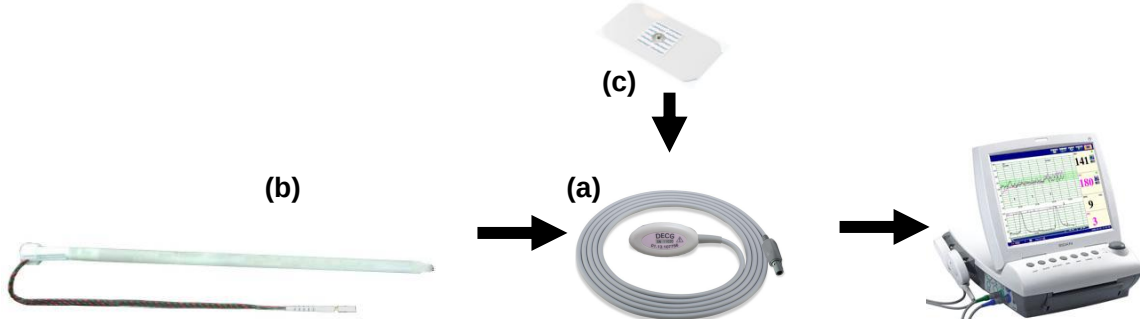


Figure 7-4 Connection for wired DECG Monitoring

Note:

The filter of AC power supply can be configured by selecting **System>Factory Configuration>Line Frequency>50(default)** or **60**. Please select it according to the current AC frequency.

7.3.4.2 Monitoring Procedure

- a) Perform a vaginal examination to identify the fetal presenting part.
- b) Prepare the patient's skin using the procedures described in section *7.3.2 Preparing the Patient's Skin Prior to Placing Electrodes*.
- c) Attach the fetal spiral electrode to the fetal presenting part using the procedures described in

section 7.3.3 *Directions for Using Fetal Spiral Electrode.*

- d) Fix an attachment pad electrode to DECG cable.
- e) Remove the film on the back of the electrode and place the electrode on maternal thigh; press it firmly in place.
- f) Connect the fetal spiral electrode to the DECG cable.
- g) Insert connector of DECG cable into the DECG socket of the monitor.

WARNING

- 1 Do not plug the fetal spiral electrode wire into the power socket.
 - 2 Never attempt to connect the fetal spiral electrode to anything other than the correct DECG cable.
-

CAUTION

Do not mistake the higher maternal heart rate for DECG.

NOTE:

- 1 If there is any doubt as to the presence of a fetal heart signal with ECG, check with the US transducer on the patient's abdomen or with a separate diagnostic instrument. The presence of an audible heart sound at a rate distinct from that of the maternal pulse is unequivocal evidence of the fetal life.
- 2 After the electrode is well attached, allow a few minutes for the electrode and fetal tissue to become stabilized. It is essential that the ECG signal electrode is in good contact with the fetal presenting part.

7.3.5 Wireless DECG Monitoring

7.3.5.1 Parts Required

- a) TOCO-E transducer
- b) FTS-3 DECG cable
- c) Fetal spiral electrode
- d) Disposable maternal attachment pad electrode

The following illustration shows how these parts should be connected:

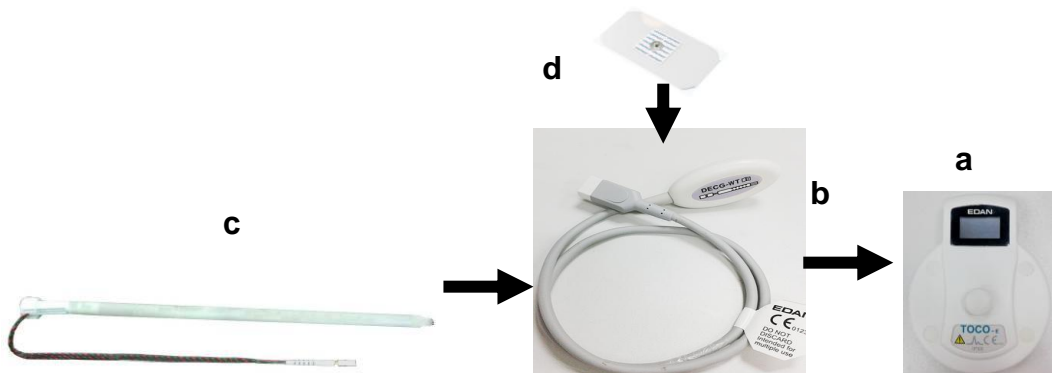


Figure 7-5 Connection for wireless DECG Monitoring

7.3.5.2 Monitoring Procedure

- a) Connect the FTS-3 system to the monitor and switch on. Refer to section 3.2.7 *connecting to the Monitor*.
- b) Take up the TOCO-E transducer and connect the FTS-3 DECG cable to it. Make sure the transducer is successfully connected to the base station.
- c) Perform a vaginal examination to identify the fetal presenting part.
- d) Prepare the patient's skin using the procedures described in section 7.3.2 *Preparing the Patient's Skin Prior to Placing Electrodes*.
- e) Attach the fetal spiral electrode to the fetal presenting part using the procedures described in section 7.3.3 *Directions for Using Fetal Spiral Electrode*.
- f) Fix an attachment pad electrode to DECG cable.
- g) Remove the film on the back of the electrode and place the electrode on maternal thigh; press it firmly in place.
- h) Connect the fetal spiral electrode to the DECG cable.

7.3.6 Changing DECG Beep Volume

When the DECG beep is enabled, the monitor gives a beep sound of DECG. The frequency of DECG beep corresponds to the fetal heart rate, but occasionally it may differ due to weak DECG signal.

To change the DECG beep volume,

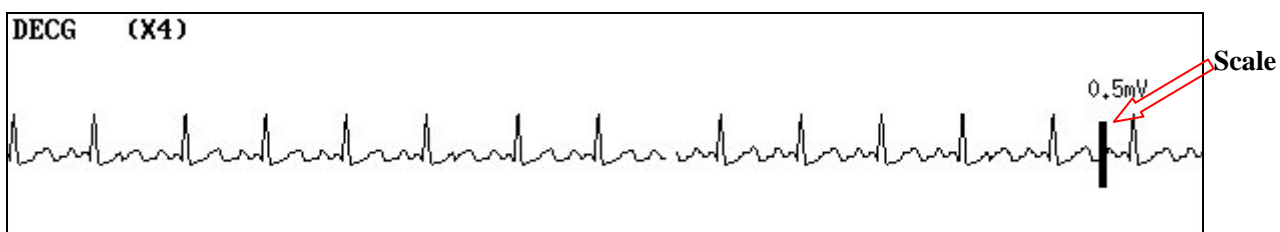
- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > DECG Beep**.
- 3 Select **0** (default) ~ **9**.
- 4 Select **OK**.

NOTE:

- 1 The DECG beep and HR beep share the same audio channel. Once the DECG beep is switched on, the HR beep is disabled (set to **OFF**) automatically.
- 2 Once the DECG/HR beep volume is changed, the sound switches to channel 1 automatically. Therefore, it is advised against changing DECG/HR beep volume in the monitoring process.

7.3.7 DECG Gain and Display

You can change the DECG gain by selecting from X1/2, X1, X2, X4 and X8 to adjust the DECG waveform size for better observation. The system displays a 0.5mV scale at the right side of the DECG waveform. The height of the 0.5mV bar is directly proportional to the waveform amplitude.



7.3.8 Switching the Artifact Suppression On or Off

When monitoring FHR with DECG, artifacts may occur due to bad connection of the spiral electrode, excessive motion of the mother, electromyographic interference etc.. The **Artifact Suppression** feature is designed to eliminate the interference. When artifact suppression is on, artifacts are suppressed and not recorded. When it is off, the artifacts are shown as well as the fetal heartbeats.

You can choose to switch the artifact suppression on or off.

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > Artifact Suppression**.
- 3 Select **ON** (default) or **OFF**.
- 4 Select **OK**.

WARNING

When artifact suppression is on, fetal arrhythmia will also be suppressed. Therefore, if fetal arrhythmia is suspected, switch artifact suppression off.

7.3.9 Detaching the Fetal Spiral Electrode

To detach the fetal spiral electrode, rotate it counterclockwise until it is free from the fetal presenting part. Do not pull the electrode from the fetal skin forcefully.

Dispose of the used fetal spiral electrode in a proper way. Do not use it again.

7.4 Monitoring Twin FHRs

7.4.1 Monitoring Twins Externally

To monitor twin FHRs externally, you need to connect a US transducer to US1 socket and the second US transducer to US2 socket of the monitor. Follow the instructions described in Section 7.2 *Monitoring FHR with Ultrasound* to acquire FHR signals for both channels. Press **CHANNEL** key to switch the FH sound from one channel to the other.

When the two US transducers are fixed, make sure FH sounds from both channels are clear, two FHR traces and two FHR numerics are displayed on the screen.

NOTE:

To ensure that both transducers stay at the optimum location, each transducer should be fixed with a separate belt.

7.4.2 Monitoring Internally

Alternatively, you can monitor a FH using ultrasound externally, and monitor the second FH using DECG internally.

Connect the US transducer to US2 socket; connect DECG cable to DECG socket.

Monitor one twin with a US transducer using the procedures described in Section 7.2 *Monitoring FHR with Ultrasound*.

Monitor the other twin with a DECG cable using the procedures described in Section 7.3 *Monitoring FHR with DECG*.

CAUTION

The US transducer must be connected to US2 socket. If the US transducer connects to US1 socket while DECG cable is connected to DECG socket, the FHR trace and numeric from US1 will not be displayed.

7.4.3 Signals Overlap Verification (SOV)

When monitoring twins, there are possibilities that one twin's FHR signal is mistaken for the other one's signal. The monitor provides signals overlap verification (SOV) function to reduce these possibilities.

In the process of monitoring, if the SOV detects signals overlapping, an alarm message "Signals Overlap (FHR1/DFHR, FHR2)" will appear on the screen to warn you. Checking the patient and reposition of transducers might be needed.

7.4.4 Changing FHR2 Offset

In order to distinguish FHR1 trace from FHR2 trace, FHR2 offset is provided to help you separate the two traces by an offset of -20 bpm or +20 bpm.

To change the FHR2 offset,

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > FHR2 Offset**.
- 3 Select **-20 bpm** (default), **0 bpm** or **+20bpm**.
- 4 Select **OK**.

This preset FHR2 offset will be printed on the recorder paper every 10 minutes (Default) /20min/30min/60min(optional) .

“FHR2: -20bpm”: the FHR2 trace is 20bpm lower than it really is.

“FHR2: 0bpm”: the FHR2 trace is in its real position.

“FHR2: 20bpm”: the FHR2 trace is 20bpm higher than it really is.

7.5 Monitoring Uterine Activity Externally

WARNING

- 1 Inspect patient skin before applying the transducer for monitoring. If the skin quality is poor, especially when the skin is damaged or irritated, please change a site to apply the transducer.
- 2 During long-time monitoring, please inspect the application site (between contractions) of TOCO transducer at least every half an hour. If the skin quality changes, you should move the transducer to another site.

7.5.1 Wired TOCO Monitoring

7.5.1.1 Parts Required

- a) TOCO transducer
- b) Belt

7.5.1.2 Monitoring Procedure

a) Placing Transducer Belt

Place the transducer belts across the bed, ensuring that the belt will be around the abdomen when it is fastened. Lay the patient on the bed.

Alternatively, the patient can take a sitting position. Arrange the belt around her abdomen.

b) Fixing the Transducer

Wipe any gel remaining on abdomen around the fundus area.

Place the TOCO transducer on the patient's abdomen, which is flat and approximately 3 cm away from the fundus, e.g. slightly above the umbilicus on the left or on the right. The

position should be different for different purposes: place the transducer close to the fetal buttocks for NST, and place it on fetal back in delivery.

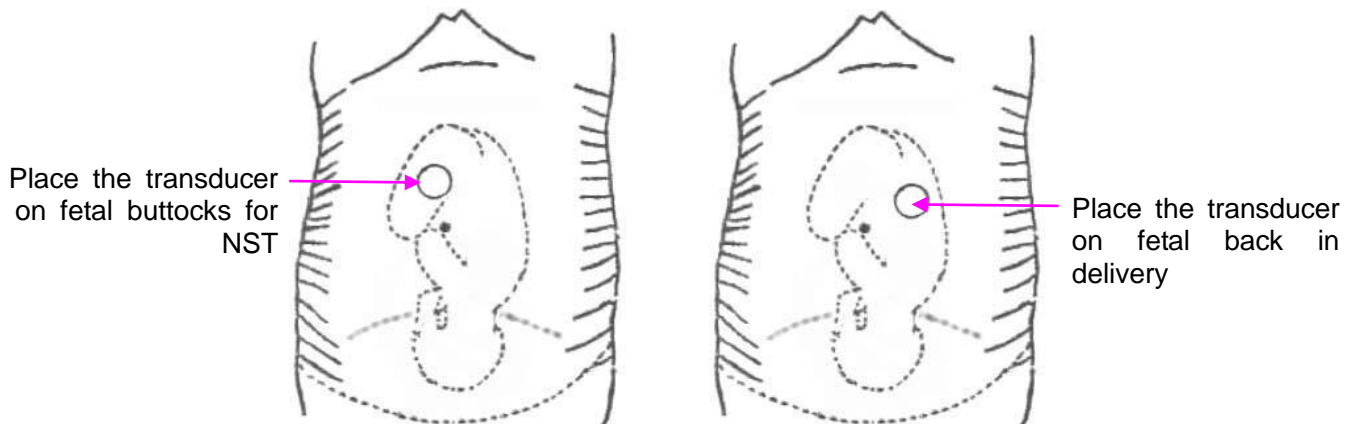


Figure 7-6 Positioning TOCO Transducer

Wrap the abdomen with the belt over the transducer. Fix the transducer by pushing its buckle through the overlapping section of the belt. Make sure the belt fits the patient snugly but comfortably.

c) *Adjusting the Numeric to Zero

Press the **AUTO ZERO** key to adjust the numeric to the baseline. Make sure this is not done during a contraction.

Wipe off any gel presents on abdomen around this area.

NOTE:

- 1 Do not apply aquasonic coupling gel on a TOCO transducer or its contact area.
- 2 Check the function of the TOCO transducer by applying pressure on it to see if this is displayed on the screen.

7.5.2 Wireless TOCO Monitoring

7.5.2.1 Parts Required

- a) TOCO-T transducer or TOCO-E transducer
- b) Belt

7.5.2.2 Monitoring Procedure

- a) Connect the FTS-3 system to the monitor and switch on. Refer to section 3.2.7 *connecting to the Monitor*.
- b) Take up the TOCO-T transducer or the TOCO-E transducer. Make sure it is successfully connected to the base station.
- c) Monitor TOCO following the procedures described in section 7.5.1.2 *Monitoring Procedure*.

7.5.3 Changing the UA Baseline

You can change the UA baseline,

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > UA Baseline**.
- 3 Select **5, 10** (default), **15** or **20**.
- 4 Select **OK**.

NOTE:

If the monitor has been configured with IUP, the IUP baseline is 0 and it is not adjustable. The TOCO baseline is adjustable.

7.5.4 Testing TOCO Transducers

To test a TOCO transducer:

- 1 Switch on the monitor.
- 2 Connect the TOCO transducer to the fetal monitor.
- 3 Gently press the center of the transducer.



Figure 7-7 Testing a TOCO Transducer

- 4 Check that the value on the display shows this change in pressure.

If a TOCO transducer fails the test, repeat this test with another transducer. If the second one passes the test, defect of the first transducer is confirmed. Replace it with a good one. If the second transducer fails the test as well, contact the manufacturer for service.

7.6 Monitoring Uterine Activity Internally

7.6.1 Parts Required

- a) Disposable intrauterine pressure catheter ACCU-TRACE™ IUPC (“IUPC” for short)
- b) Reusable intrauterine pressure cable (“IUP cable” for short)
- c) Reusable intrauterine pressure connecting cable (“connecting cable” for short)

The following illustration shows how these parts should be connected:

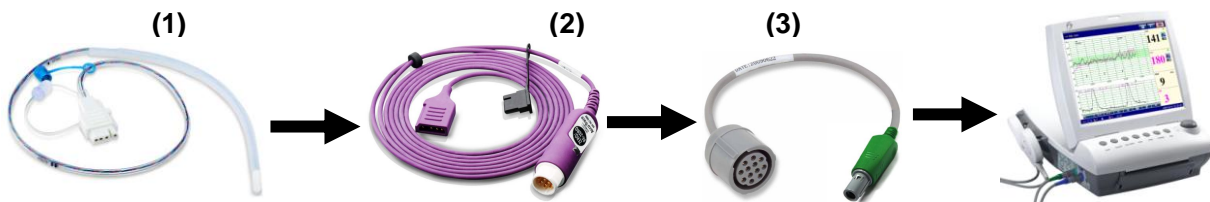


Figure 7-8 Connection for IUP Monitoring

7.6.2 Directions for Use of IUPC

Preparation

- 1) Gather supplies: ACCU-TRACE IUPC, reusable cable, and amnioinfusion supplies if needed.
- 2) Open the sterile ACCU-TRACE IUPC package.

Insertion

NOTE:

This product is designed for use with the introducer.

- 3) Using aseptic technique, remove the catheter from the package.
- 4) Perform vaginal exam to ensure ruptured membranes and adequate dilation.
- 5) Advance the catheter tip to the cervical os along the examination hand, using the hand as a guide. Do not advance the introducer through the cervix.
- 6) Continue to gently advance the catheter tip through the cervical os and feed the catheter into the intra-amniotic cavity until the 45 cm mark is at the introitus. If the 45cm mark is not clearly visible, stop advancing when the ● symbol on the catheter meets the introducer.

NOTE:

For easier insertion, do not twist the catheter in the introducer.

- 7) The IUPC may be spontaneously filled with amniotic fluid. This can be seen in the clear lumen of the catheter. The filter cap will prevent the amniotic fluid from leaking.
- 8) Slide the introducer out of the vagina along the catheter. When the introducer is

completely out of the vagina, slide thumb between catheter and introducer tab, which will begin to separate the introducer from the catheter.

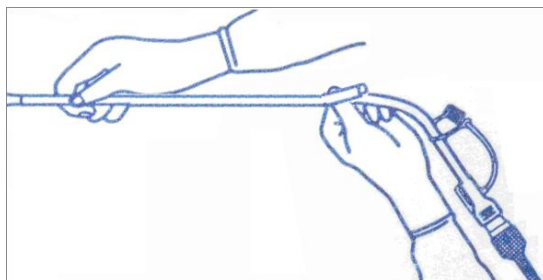


Figure 7-9 Separate the Introducer

- 9) Anchor the catheter in place with one hand, and pull the introducer straight back off the catheter.

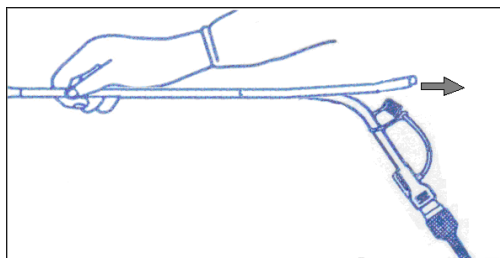


Figure 7-10 Remove the Introducer

- 10) Remove the liner from the adhesive pad, and then adhere the pad to the patient's skin. Secure the catheter by placing the catheter attachment strap to the adhesive pad..

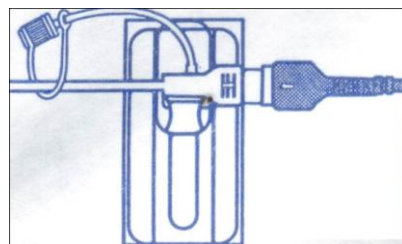


Figure 7-11 Secure the Adhesive Pad to Mother

Rezeroing the System During Monitoring

- 1) With the catheter connected to the IUP cable, momentarily pressing the re-zero button on the pressure cable. The green light on the cable will flash for five seconds.

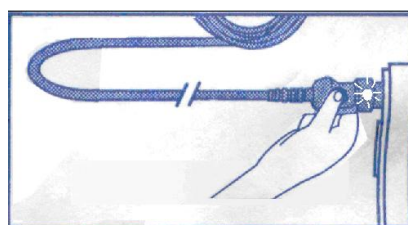


Figure 7-12 Rezeroing the System

- 2) During this period, adjust the monitor to zero by pressing **AUTO ZERO** key.

WARNING

- 1 Before insertion of IUPC, placental position should be confirmed, amniotic membranes are adequately ruptured and sufficient cervical dilatation is assured.
- 2 Try to insert the catheter opposite the placental site. Do not insert the introducer beyond the cervical OS. Use it with caution when uterine infection is present.
- 3 If resistance is met at any time during insertion, withdraw the catheter slightly and try at a different angle. Forced insertion may result in patient's discomfort or injury.

CAUTION

- 1 Since procedures vary according to hospital needs/preferences, it is the responsibility of the hospital staff to determine exact policies and procedures for both monitoring and amnioinfusion. The safe and effective use of the IUPC depends on the skill of the clinician who applies /uses it.
- 2 The IUPC has been sterilized by gamma radiation and is sterilized and non-pyrogenic unless package is broken or open. Do not re-sterilize it.

NOTE:

Refer to the instruction on the package for more information about using the IUPC.

7.6.3 IUP Monitoring Procedure

- 1) Insert IUPC using the procedure described in section 7.6.2 *Directions for Use of IUPC*.
- 2) Connect the IUPC to the IUP cable.

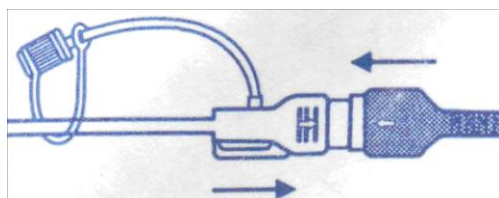


Figure 7-13 Connect Catheter to Pressure Cable

- 3) Connect the IUP cable to the connecting cable. (They might have already been well connected in the package.)
- 4) Plug the connecting cable to the TOCO/IUP socket of the monitor.
- 5) Momentarily pressing the re-zero button on the IUP cable. The green light on the cable will flash for five seconds. During this period, zero the monitor by pressing the **AUTO ZERO** key. Make sure the display numeric and trace are both “0”.
- 6) Ask the mother to cough. A spike on the trace in response to the cough indicates proper positioning and function of the IUPC.
- 7) Wash timely during monitoring. A spike on the tracing will respond to the washing.

7.6.4 Checking Intrauterine Pressure Cable Function

To test an IUP cable's function:

- 1) Disconnect the catheter from the cable. Insert the cable check plug into the catheter end of the cable.

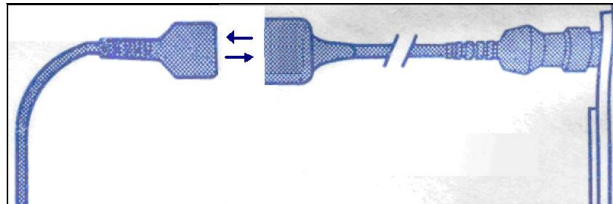


Figure 7-14 Test the Pressure Cable

- 2) Verify that the green light is continuously lit (no flashing).
- 3) If the light does not illuminate, replace the cable.

NOTE:

- 1 If the light is flashing, verify that the cable check plug is inserted completely into the cable.
- 2 The cable test function is not intended to check the accuracy of the system, only to confirm cable function.

7.7 Monitoring Fetal Movement

7.7.1 Auto Fetal Movement (AFM) Monitoring

During fetal heart monitoring with ultrasound, the fetal movement signals are also detected. The fetal movement signals differ from the Doppler heart rate signals in that they have larger extent and lower frequency. The larger extent is because of the bigger scope of moving areas (e.g., the fetal arms or legs); lower frequency is because of the lower velocity of the fetal movements compared with those of the fetal heart.

Only US1 channel can perform AFM. But be aware that when monitoring twins, the movements detected by US1 may also be caused by the second fetus's movement.

The movement of the fetus will be detected and displayed in the form of a trace on the screen and the recorder paper.

There are two ways to monitor automatic fetal movement (AFM): wired monitoring and wireless monitoring. Wired AFM monitoring is monitored by wired ultrasound transducer while wireless AFM monitoring is monitored by wireless ultrasound transducer (US-T transducer). AFM can be monitored simultaneously with FHR. Be aware that only when the wired ultrasound or the US-T transducer is connected to US1 channel can AFM be monitored.

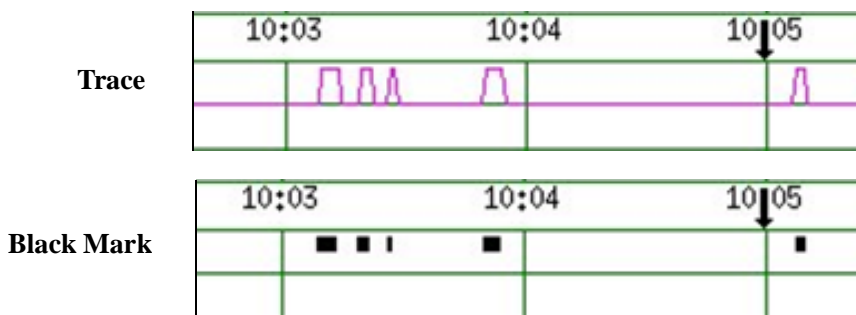
7.7.2 Enabling or Disabling AFM Monitoring

To enable or disable AFM monitoring,

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > AFM**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

7.7.3 Choosing AFM Mode

When AFM monitoring is enabled, the AFM monitoring result is displayed either in the form of a trace or black marks. The x-axis of each wave or each black mark indicates the duration of a detected fetal movement.



To choose AFM mode,

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > AFM Mode**.
- 3 Select **Trace** (default) or **Black Mark**.
- 4 Select **OK**.

7.7.4 Choosing FM Source

When AFM monitoring is enabled, the FM has two sources: AFM and MFM.

To choose the FM source,

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > FM Source**.
- 3 Select **MFM** (default) or **AFM**.
- 4 Select **OK**.

7.7.5 Manual Fetal Movement (MFM) Monitoring

MFM result comes from the patient's feeling of fetal movement. The count will be displayed on

the screen in MFM numeric area.

- 1) Insert the FM marker connector into the **MARK** socket on the monitor.
- 2) Let the patient hold the marker in hand; ask her to press the top key of it when a fetal movement is felt. Continuous movements in 5 seconds are considered to be one movement and only press the key once.

7.7.6 Changing MFM Volume

The monitor gives a sound when the FM marker key is pressed, and the volume is adjustable.

To change the MFM volume,

- 1 Select the setup key  on the main interface.
- 2 Select **Fetus > MFM Volume**.
- 3 Select **Low** or **High** (default).
- 4 Select **OK**.

7.8 *Start Monitoring

After the **START** key is pressed, the monitor automatically zeroes the pressure, clears the FM count and starts monitoring.

If the Auto start printing is disabled, press the **PRINT** key to start printing.

7.9 *Inputting Maternal Information (Mat. Info)

7.9.1 Auto ID

After you press the **START** key, the system creates an auto-ID for the present patient. (if Mat. Info inputting is switched off.) The auto-ID consists of the date and time when the monitoring starts.

7.9.2 Changing Maternal Information

You can change the patient's information after the monitoring starts:

- 1 Select Mat. Info key  on the main interface.

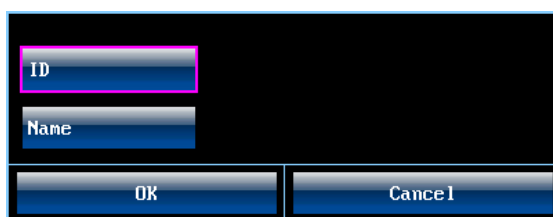


Figure 7-15 Mat. Info Inputting Menu

2 Select ID.

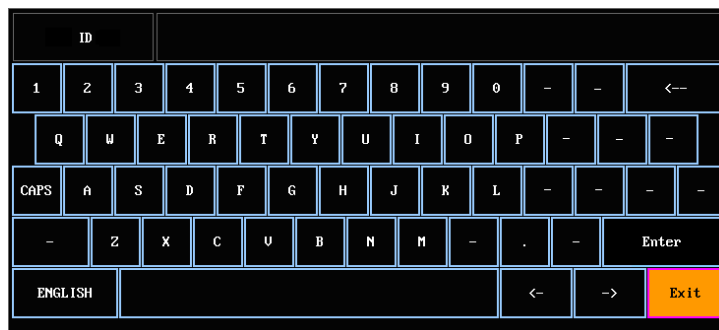


Figure 7-16 Soft Keyboard

- 3 Select the required character for patient's ID on the soft keyboard.
- 4 Select **Enter**.
- 5 Select **Name**.
- 6 Select the required letter for patient's name on the soft keyboard.
- 7 Select **Enter**.
- 8 Select **OK**.

The monitoring does not stop when you change maternal information. After you select **OK** to exit, the new ID takes the place of the old one for this patient.

NOTE:

- 1 Pressing the **START** key separates two patients. The monitor only displays the most recent ID for the same patient.
- 2 If printing starts automatically with the monitoring, the first ID printed on the recorder paper will be the auto-ID. The new ID will be printed 10 minutes/20 minutes/30 minutes/ 60 minutes(optional) later.
- 3 The ID and name are shown on the screen, the paper printout and the archive list.
- 4 For the non-English system, more letters are provided for inputting the name. Select the key on the bottom left corner to toggle between them.
- 5 When the network version is ETHERNET 1.4, if ID is revised on either the fetal monitor or the central station after confirmation, they will synchronize ID with each other.

7.9.3 Switching Mat. Info Inputting On or Off

The **Mat. Info inputting** function allows the menu to pop up automatically after the **START** key is pressed. After you input the mother's information and exit from the menu, the monitoring starts immediately.

To switch the **Mat. Info Inputting** on or off:

- 1 Select the setup key  on the main interface.
- 2 Select **Start Monitoring > Mat. Info.**
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

Chapter 8 Fetal Monitoring Display (F9)

8.1 Traces

WARNING

- 1 Due to the LCD size, resolution and system settings, the traces displayed on the screen may look different from the recorder printout. The printout should prevail when making diagnoses.
- 2 If the data is doubtful, clinicians should make diagnoses based on the real condition.

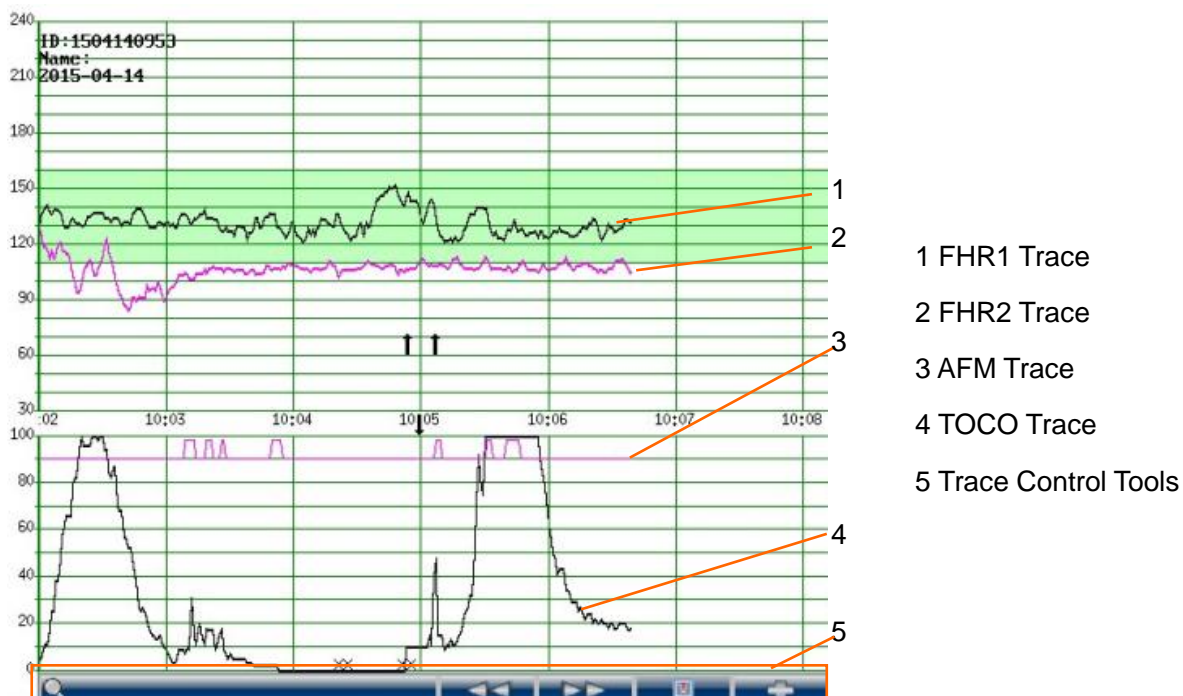


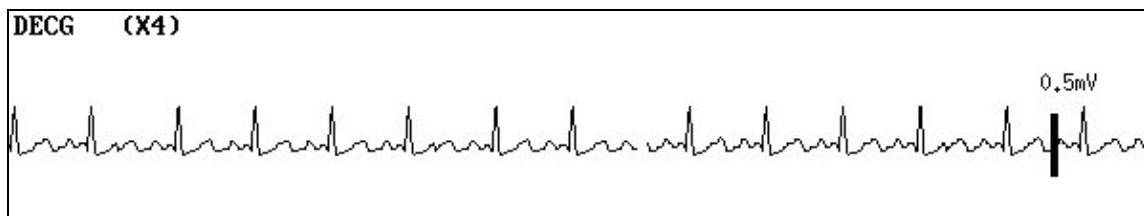
Figure 8-1 Traces

During monitoring or reviewing, the trace window displays four traces: FHR1 trace, FHR2 trace (dual configuration), AFM trace and TOCO trace.

FHR1/FHR2 trace

The y-axis of the trace indicates the numerics of FHR. The range is 30 bpm ~ 240 bpm (American standard) or 50 bpm ~ 210 bpm (International standard).

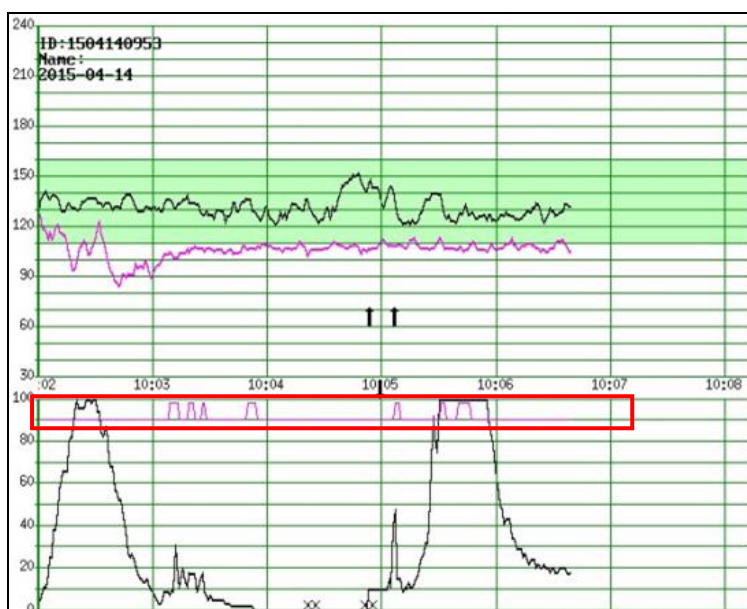
If FHR is monitored using DECG, and the DECG trace is switched on in the hardware setup (only service engineers have access to it), a DECG trace is shown underneath other traces on the screen.



The AFM monitoring result is displayed either in the form of a trace or in the form of black marks.

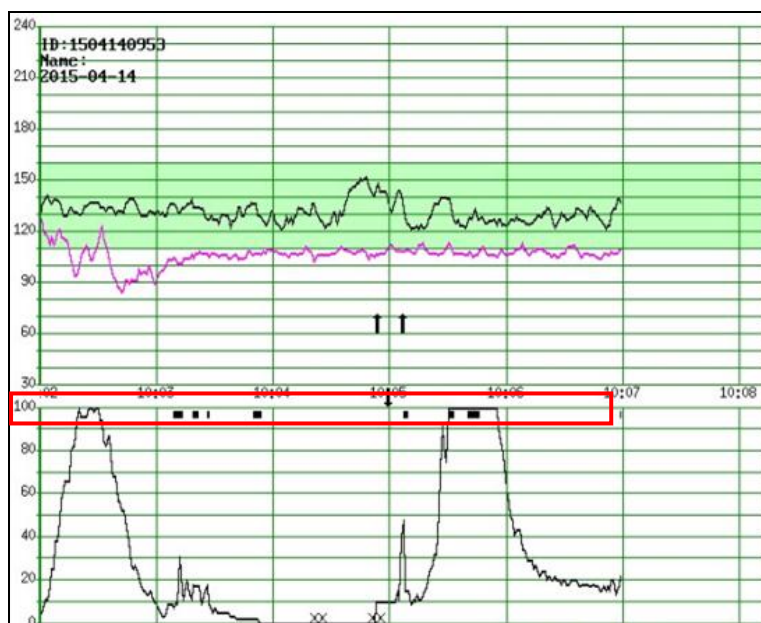
AFM trace

The x-axis of each wave indicates the duration of a detected fetal movement.



AFM black mark

The x-axis of each black mark indicates the duration of a detected fetal movement.



NOTE:

AFM trace is only for reference. Please take the MFM marks as criterion.

TOCO trace

The y-axis indicates the numeric of TOCO. The range is 0 ~ 100.

Besides, some other symbols appear among the traces:

- This symbol indicates the new monitoring starts.
- ↑ This symbol indicates a manual fetal movement, and it appears after the patient presses the FM marker when she feels a fetal movement.
- ↓ This symbol indicates the **MARK** key is pressed to record an event, such as the patient turning around, taking injection.
- ☒ This symbol indicates the monitor is zeroed by pressing **AUTO ZERO** key.

8.1.1 Changing Time Scale

The fetal monitoring traces share the same time scale. This scale is either in real time format or relative time format. Real time is the time of the monitor. Relative time records the elapsed time for the current monitoring.

To change this time format:

- 1 Select the setup key  on the main interface.
- 2 Select **Date And Time > Time Scale**.
- 3 Select **Real Time** (default) or **Relative Time**.
- 4 Select **OK**.

NOTE:

The real time contains only the hour and minute, but no second. As a result, the time scale may correspond to the 0 ~ 59th second of the system time. Do not mistake the time scale for the exact time.

8.2 Trace Control Tools

Figure 8-2 Trace Control Tools

- | | |
|-----------------------|---------------------|
| 1 Searching Key | 2 Reviewing Keys |
| 3 Alarm Reviewing Key | 4 CTG Analyzing Key |

8.2.1 Data Saving

When the **START** key is pressed, the monitor saves data of the previous ID in a file, and then clears it from the main interface. The main interface only displays the new patient's data. During monitoring, the data is saved every 10 minutes. All data of the same patient is saved in a file (the maximum duration is 24 hours, the rest data is saved in another file.)

The files are stored in the monitor. When the data amount reaches the maximum capacity (300 files or approximately 60-hour), the monitor deletes the oldest file(s) automatically.

When the network version is ETHERNET 1.4, if the central station disconnects with the fetal monitor in one minute, the monitoring data will be re-uploaded to the central station after re-connection, if the disconnection is over one minute, then the monitoring data can't be uploaded to the central station.

8.2.2 *Searching for a File

When the USB facility is disabled, the searching key  under the traces is used to search for a patient's data file saved in the monitor.

To search for a patient,

- 1 Select the searching key to open the file list. It contains six sets of most recent patients' ID, name and start time of monitoring. Select the required item, this file is loaded to the main interface immediately.



Figure 8-3 File List

If the required file is not in this list,

- 2 Select **MORE** to open the **Patient Searching** window.

<< Patient Searching >>		
ID	Name	Search
ID	Date	Name
888888	2015-03-18 14:35:30	IIII
777777	2015-03-18 14:31:14	UUUUU
666666	2015-03-18 14:19:38	YYYYY
555555	2015-03-18 14:14:40	TTTTTT
444444	2015-03-18 14:10:02	RRRRR
333333	2015-03-18 14:00:42	EEEEEE
222222	2015-03-18 13:48:57	WWWWW
111111	2015-03-18 13:47:34	QQQQQ

Cancel

Figure 8-4 Patient Searching

3 Select **ID**, input the patient ID with the soft keyboard and select **Enter**.

4 Select **Name**, input the patient's name with the soft keyboard and select **Enter**.

NOTE:

You can input only a part of the patient ID or name. However, the more information you input, the more accurate result you will get.

5 Select **Search**. The files with the matched information are listed in the window.

6 Select the required item, and this file is loaded to the main interface immediately. You can review the traces backward or forward.

8.2.3 File Management (Optional)

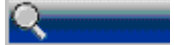
The USB facility of the monitor allows you to export the auto-saved files into a USB disk, and then you can save the files in a PC or open them in a data managing system.

Once the monitor is configured with the relevant hardware, the USB feature can be enabled or disable by the service personnel of the manufacturer.

8.2.3.1 Exporting Files

1 Make sure the USB feature is enabled. Stop printing and disconnect the network.

2 Plug the USB disk into the USB socket on rear panel of the monitor (figure 2-4). A message "Ready to use USB disk" in the message area indicates the proper insertion of the disk.

3 Select the file managing key  on the main interface to open the File Management interface, which records a list of up to 300 most recent monitoring records (patients' ID, name and date) and a few operation items.

<< File Management >>		
ID	Name	Search
ID	Date	Name
1709041559	2017-09-04 15:59:40	Lucy
1708251120	2017-08-25 11:20:56	Lily
1708251117	2017-08-25 11:17:38	Katty
1708241501	2017-08-24 15:01:50	Aimee
1708231320	2017-08-23 13:20:45	Anne
1708221730	2017-08-22 17:30:46	Kerry
1708211257	2017-08-21 12:57:37	Alice
1708161532	2017-08-16 15:32:53	Ailsa

Figure 8-6 File Management

- 4 If the required record is not on the current page, select Next to view more records.
- 5 Move the cursor to select the required item, and then select Export in the pop-up item, and the monitor exports this record to the USB disk. Or you can select Export All to export all the records to the USB disk.

NOTE:

- 1 When the monitor is in the process of printing or is connected to the network, the files cannot be exported.
- 2 To avoid influence caused to the real-time monitoring, the manufacturer advises against plugging in the USB disk and exporting the data in the process of monitoring.
- 3 The USB disk is not a tool for long-term data storage. You should save the exported files in a PC in time and clear the USB disk termly.
- 4 The monitor only supports those USB disks with **FAT** or **FAT32** (recommended) format, and with capacity not larger than 8G. You are advised to use the USB disk provided by the manufacturer.

In the **FetusData** folder of the USB disk, a sub-folder named after the export date and time is created when the export is performed. The exported records are saved in this sub-folder as .trc files, named after the monitor started date, time and ID, e.g. “20100120-124936-12345.trc”.

8.2.3.2 Removing USB Disk

After the export finishes, select **Remove Disk** on the **File Management** interface. Do not unplug the USB disk until a message “The USB can now be safely removed.” is prompted.

If the message “Failure” is prompted, you should perform the above procedures again.

NOTE:

- 1 Make sure you perform the **Remove Disk** procedure, otherwise data lose or USB disk

damage may be caused.

- 2 You should unplug the USB disk after performing the **Remove Disk** procedure; otherwise the monitor cannot identify the USB disk.

8.2.3.3 Deleting Files

After the files are saved, you can delete them.

1 Select the file managing key  on the main interface to open the **File Management** interface.

2 Select Delete All > Yes. All the files in the monitor are deleted.

3 Select Exit.

NOTE:

- 1 When the monitor is in the process of printing, the files cannot be deleted.
- 2 File deleting should be performed with caution since the deleted files cannot be restored.
- 3 The monitor automatically erases the earliest files when the memory is full (the maximum capacity is 60-hour data). You should export and save the files in time.
- 4 When there are more than 300 files, it may take extended time for the monitor to load them. You should export the files in time and then delete them from the monitor.

8.2.4 *Reviewing

The reviewing keys  (backward key) and  (forward key) are used to review the traces. The word **REVIEW** is shown in the background when reviewing the traces.

Select the backward key to review the previous traces. The traces start to retreat. The amount of the progress symbol “<” on top of the traces indicates the retreating speed. Rotate the control knob anticlockwise or touch the  symbol to increase the speed until it reaches the maximum.

Rotate the knob clockwise or touch the  symbol to decrease the speed until it reaches the minimum. Press the knob or touch any place on the screen to pause.

Select the forward key to review the next traces. The traces start to advance. The amount of the progress symbol “>” on top of the traces indicates the advancing speed. Rotate the control knob clockwise or touch the  symbol to increase the speed until it reaches the maximum. Rotate the knob anticlockwise or touch the  symbol to decrease the speed until it reaches the minimum. Press the knob or touch any place on the screen to pause.

When the reviewing is paused, the progress symbol turns to <--X%-->. If the **PRINT** key is pressed at this moment, the recorder will print the traces starting from the left edge of the screen at a high speed.

X% indicates the proportion of current traces positioned in the whole reviewable traces.

Move the cursor away from the trace control tools, or touch any place out of the trace window on the screen to return to the real-time main interface. If no operation is performed in 10 seconds, the monitor switches to real-time interface automatically, unless the printing is in process.

When reviewing the traces, the monitor does not stop. The FH sound and numerics are all real time information of the current patient.

WARNING

The reviewing printout is provided for reference only. Please take the real-time printout as criterion when making diagnoses.

NOTE:

- 1 The main interface only displays traces and patient information of one file. If you want to review another file you should search for the file and load it.
- 2 For a real-time monitoring patient, you can print all her traces, including SpO₂ trace. However, when printing traces in a file, the SpO₂ trace cannot be printed.
- 3 You must pause before printing starts. Printing in the process of playback might result in failure information on the paper.
- 4 After the reviewed data has been printed out, the recorder does not switch back to real-time printing automatically.

8.2.5 *CTG Analysis

CTG analysis aims at a real-time trace, providing some reference data for the physicians. It only analyzes the real-time trace after it's been printed for 10 minutes, and the longest duration is 60 minutes.

WARNING

- 1 CTG analysis is used for the surveillance of pregnancies and not in delivery room of childbirth.
- 2 CTG analysis is just an analysis intended to assist the physicians in interpreting the waveforms. Conclusions should be drawn on the basis of the physicians' diagnosis.
- 3 This analysis describes the fetal heart rate, the tocography and the fetal movements. It's the responsibility of qualified medical staff to do the diagnostic interpretation of the waveform.

8.2.5.1 Enabling/Disabling CTG Analysis

- 1 Select the setup key  on the main interface.
- 2 Select **General > CTG Analysis**.
- 3 Select **ON** or **OFF** (default).

4 Select **OK**.

A CTG analysis key  appears on the main interface, indicating that CTG analysis is enabled.

8.2.5.2 CTG Analyzing

NOTE:

- 1 CTG analyze starts after the real-time trace has been printed for 10 minutes.
- 2 The CTG analysis result is for reference only.

After the real-time trace is printed for 10 minutes, select the CTG analysis key  on the main interface. The analysis result window opens.

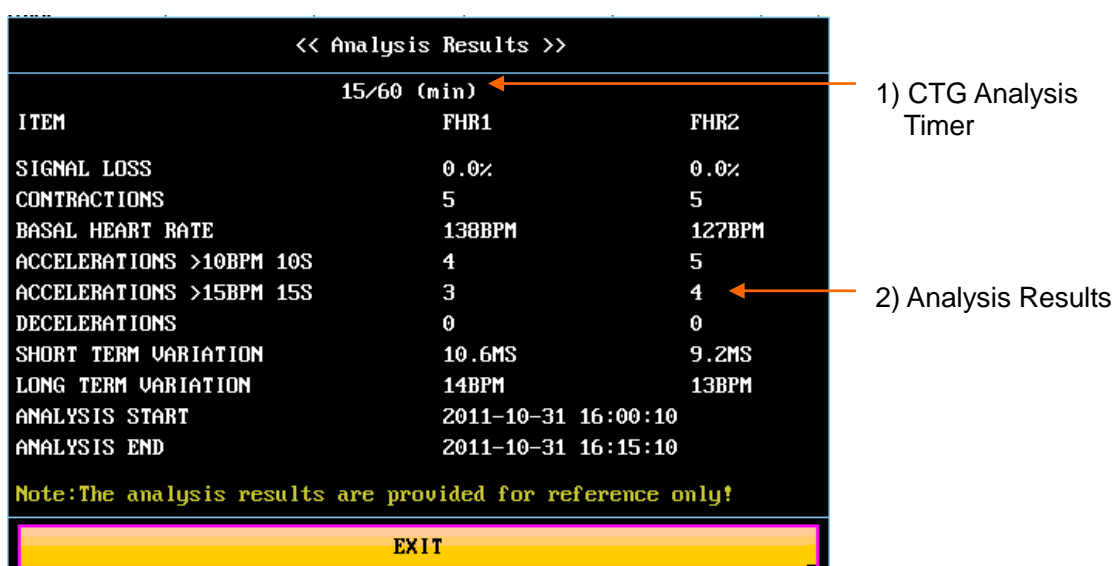


Figure 8-5 CTG Analysis Results

Refer to figure 8-5, the CTG analysis results on the screen include:

1) CTG Analysis Timer:

The CTG analysis timer starts when the recorder starts printing; it stops when the timer reaches 60 minutes (the timer turns into **>60**) and resets when the recorder stops printing.

2) CTG Analysis Results:

SIGNAL LOSS:	the proportion of the signal loss. If it is larger than 10%, analysis results cannot be acquired.
CONTRACTIONS:	the contraction time during analysis.
BASAL HEART RATE:	the average FHR in 10 minutes when it is not influenced by fetal movement or contractions.
ACCELERATIONS:	the acceleration time, including the acceleration with amplitude larger than 10bpm and lasts more than 10 seconds, and the acceleration with amplitude larger than 15bpm and lasts more than

	15 seconds.
DECELERATIONS:	the deceleration time.
SHORT TERM VARIATION:	the short-term variation analysis result.
LONG TERM VARIATION:	the long-term variation analysis result.
ANALYSIS START:	the start time of the analysis.
ANALYSIS END:	the finishing time of the analysis.

During 10 to 60-minute of the timer, the monitor gives CTG analysis results every minute.

At the end of the printing, the recorder prints the CTG analysis results of this moment on the recorder paper.

Be aware that CTG analysis result is a calculation output. It can be used as a reference to assist medical personnel in making correct diagnosis, instead of replacing it.

NOTE:

Do not disconnect the ultrasound transducer(s) before the printing stops, otherwise the analysis results will not be printed.

8.2.6 *Marking a Note

When there is a significant event, you can press the **MARK** key on the front panel to add a note. An event mark ↓ will appear on both the main interface and the recorder paper.

However, an event mark cannot clearly indicate an event. **Smart Notes** provides a list of annotation for the events, including events that relate to drugs, positions, membranes, procedures, antenatal, reasons and others.

8.2.6.1 Enabling/Disabling Smart Notes

To enable or disable **Smart Notes**,

- 1 Select the setup key  on the main interface.
- 2 Select **General > Smart Notes**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

A smart note editing key  appears next to the **Smart Notes** item.

8.2.6.2 Annotating an event

Once **Smart Notes** is enabled, press the **MARK** key on the front panel to open the smart note list, choose an event catalog and then choose an annotation from the list.

The annotation of this event will be printed in the top area of recorder paper during real-time printing.

8.2.6.3 Changing smart note content

You can change the smart note content in the smart note list by performing the following steps:

- 1 Select the setup key  on the main interface.
- 2 Select **General**.
- 3 Select the smart note editing key .
- 4 Select a catalog.
- 5 Select a note.
- 6 Use the soft keyboard to edit the note content.
- 7 Select **Enter**.
- 8 Select **OK**.

8.3 Numerics

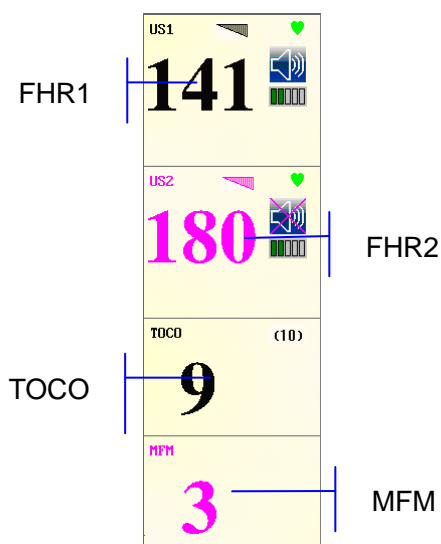
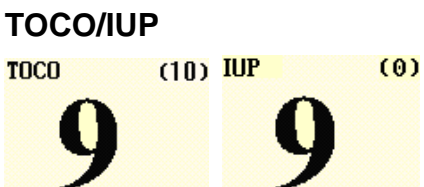


Figure 8-6 Fetal Monitoring Numerics

The fetal monitoring values in the numeric window include FHR1/DFHR value, FHR2 value, TOCO/IUP value and FM count:

FHR1/DFHR		 : FHR signal quality. When the quality is poor, it turns into  .
		 : FH refreshing rate
		 : FH sound volume adjusting key.
		 : FH sound volume indicator

	141: FHR1/DFHR measurement numeric. If the US1 socket is not connected with a US transducer and DECG socket is not connected with a DECG cable, nothing displays here; if the transducer/cable is connected but no monitoring is going on, it displays ---.
	180: FHR2 measurement numeric. If the US2 socket is not connected with a US transducer when switching on, it displays OFF but no numeric here; if the transducer is connected but no monitoring is going on, it displays ---.
	(10)/(0): UA baseline 9: current UA measurement numeric
	MFM/AFM: FM source 3: FM count

When F9 Fetal Monitor is connected to FTS-3 Telemetry System, the signal strength icon , signal strength and quality icon , and battery level icon  of the US-T transducers and TOCO-T transducer or TOCO-E transducer are displayed in the numeric window.

8.3.1 Changing Numeric Window Position (F9)

Especially for **F9**, the numeric window can be located either on the right of the traces or on top of them. To change its position,

- 1 Select the setup key  on the main interface.
- 2 Select **General > Numeric Window**.
- 3 Select **Top** or **Right** (default).
- 4 Select **OK**.

Chapter 9 Maternal Monitoring (F9 Express)

WARNING

- 1 Do not apply this monitor during electro-surgery or MRI; otherwise it might result in harming the patient or the operator.
- 2 Always check if the alarm settings are appropriate for your patient before starting monitoring.
- 3 Check for any fault of the transducers before applying them to the patient.

NOTE:

This feature is only available on **F9 Express**.

9.1 Maternal ECG Monitoring

9.1.1 Introduction

ECG monitoring produces a continuous wave form of the patient's cardiac electric activity to enable an accurate assessment of current physiological state. Only proper connection of ECG cables can ensure a satisfactory measurement.

A 20-second monitor stabilization period shall be allowed before testing. The monitor has Tall T-wave rejection capability.

The response time of heart rate meter to change in heart rate is less than 10s.

The minute heart rate display is updated at an interval of 1s.

The monitor computes heart rate by excluding the minimum and maximum values from the 12 most recent RR intervals and averaging the residual 10 RR intervals. If each of three consecutive RR intervals is greater than 1200ms, then the four most recent RR intervals are averaged to compute the MHR.

FTS-3 system computes heart rate by averaging the 12 most recent RR intervals.

The monitor or FTS-3 system does not have capability of detecting or rejecting pacemaker pulse, nor does it provide a pulse to synchronize a defibrillator discharge.

The monitor does not give alarm for tachycardia and cardiac arrhythmia.

The d.c. offset voltage tolerance of the monitor is from -500mV to +500mV. If the d.c. offset voltage of the detected ECG signal is out of this range, the monitor issues a high level alarm: ECG SINGNAL EXCEEDS LIMIT.

WARNING

- 1 When connecting the cables and electrodes, make sure no conductive part is in contact with the ground. Verify that all ECG electrodes, including neutral electrodes, are securely attached to the patient.
- 2 The electrodes should be made of the same metal materials.

WARNING

- 3 ECG accessories are not suitable for DIRECT CARDIAC APPLICATION (Refer to IEC60601-1 for more information about the definition of DIRECT CARDIAC APPLICATION).

CAUTION

A different type of electrodes may produce higher offset voltage. Therefore, only use the ECG leads supplied by the manufacturer when using the monitor for ECG monitoring.

NOTE:

Interference from a non-grounded instrument near the patient and ESU interference can cause inaccuracy of the waveform.

A good ECG signal should be –

- 1) With normal QRS wave.
- 2) Tall and narrow with no notches.
- 3) With tall R-wave completely above the baseline.
- 4) With T-wave less than one-third of the R-wave height.
- 5) With P-wave much smaller than the T-wave.

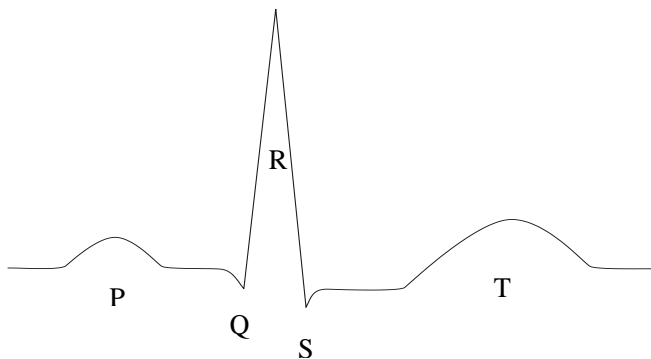


Figure 9-1 Standard ECG Waveform

9.1.2 How to Place 3-lead ECG Cables

The table below lists the names and position of 3-lead ECG cable in America and Europe.

AHA		IEC		Position
Name	Color	Name	Color	
RA	White	R	Red	Near the right shoulder, right below the clavicle
LA	Black	L	Yellow	Near the left shoulder, right below the clavicle
LL	Red	F	Green	On the left hypogastrium

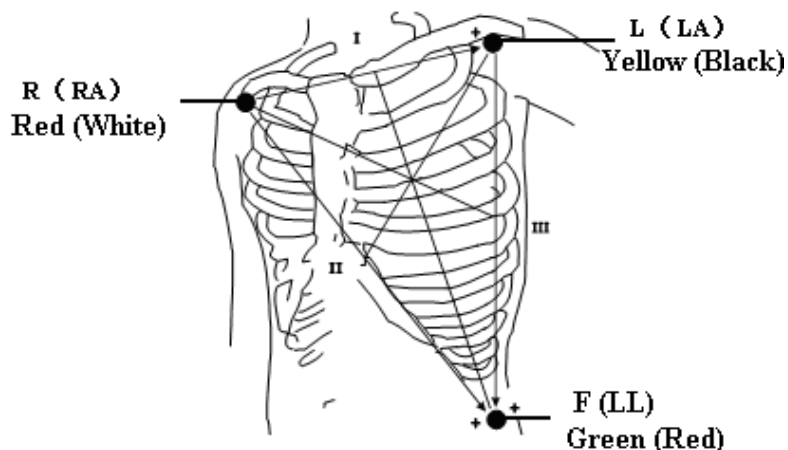


Figure 9-2 Placing 3-lead ECG Cable

NOTE:

- 1 To ensure patient's safety, all leads must be attached to the patient.
- 2 Check everyday if the skin is irritated from attachment of electrodes, if so, change for new electrodes or change their sites every 24 hours.
- 3 Recycle or dispose the used electrodes properly to protect the environment.

9.1.3 How to Place 2-lead FTS-3 MECG Cables

The table below lists the names and position of 2-lead FTS-3 MECG cable in America and Europe.

AHA		IEC		Position
Name	Color	Name	Color	
RA	White	R	Red	Near the right shoulder, right below the clavicle
LA	Black	L	Yellow	Near the left shoulder, right below the clavicle

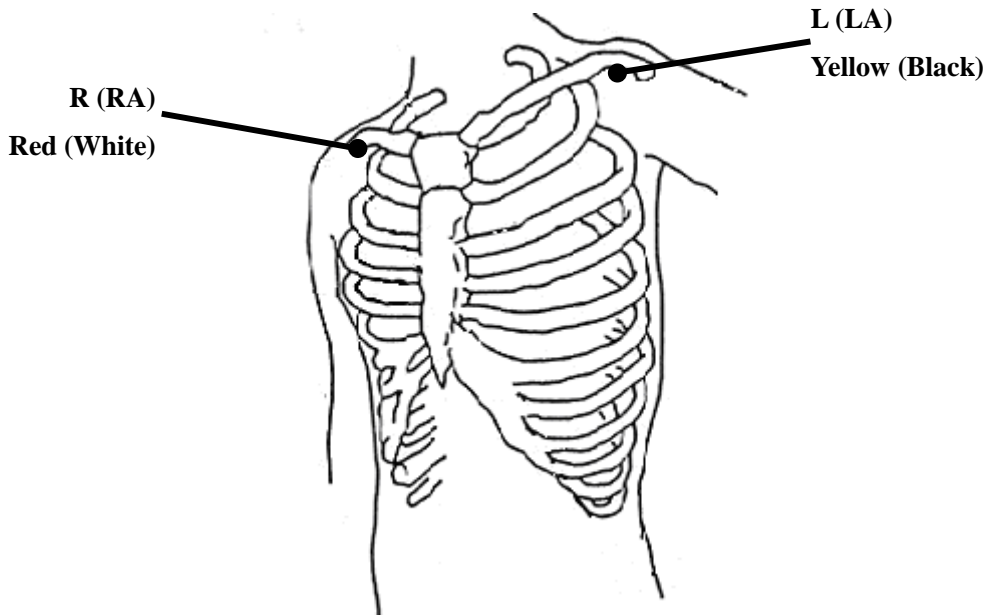


Figure 9-3 Placing 2-lead FTS-3 MECG Cable(1)

If the signal is not good, please try below position.

AHA		IEC		Position
Name	Color	Name	Color	
RA	White	R	Red	On the right side of the navel, the brim of the electrode is about 4 fingers' breadth from the navel.
LA	Black	L	Yellow	The crossing point of the left anterior axillary line and the fifth rib.

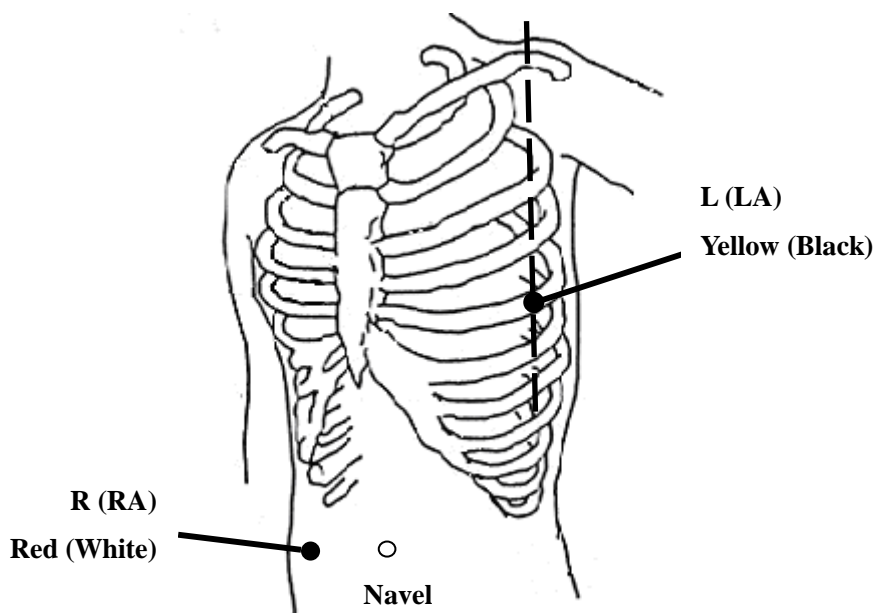


Figure 9-4 Placing 2-lead FTS-3 MECG Cable(2)

9.1.4 Wired ECG Monitoring

9.1.4.1 Parts Required

- a) 3-lead ECG cable
- b) Attachment pad electrodes

Note:

The filter of AC power supply can be configured by selecting **System>Factory Configuration>Line Frequency>50(default)** or **60**. Please select it according to the current AC frequency.

9.1.4.2 Monitoring Procedure

- a) Prepare the skin for ECG monitoring. Refer to section 7.3.2 *Preparing the Patient's Skin Prior to Placing Electrodes*.
- b) Insert the ECG cable connector into the MECG socket on the monitor.
- c) Connect attachment pad electrodes with an ECG cable.
- d) Peel the protection membrane off the back of attachment pad electrodes and attach electrodes to the patient. Refer to section 9.1.2 *How to Place 3-lead ECG Cables* for electrodes' sites.

9.1.5 Wireless ECG Monitoring

9.1.5.1 Parts Required

- a) TOCO-E transducer
- b) 2-lead FTS-3 MECG cable
- c) Attachment pad electrodes

9.1.5.2 Monitoring Procedure

- a) Connect the FTS-3 system to the monitor and switch on. Refer to section 3.2.7 *connecting to the Monitor*.
- b) Take up the TOCO-E transducer and connect the FTS-3 MECG cable to it. Make sure the transducer is successfully connected to the base station.
- c) Prepare the skin for ECG monitoring. Refer to section 7.3.2 *Preparing the Patient's Skin Prior to Placing Electrodes*.
- d) Connect attachment pad electrodes with the FTS-3 MECG cable.
- e) Peel the protection membrane off the back of attachment pad electrodes and attach electrodes to the patient. Refer to section 9.1.3 *How to Place 2-lead FTS-3 MECG Cables* for electrodes' sites.

WARNING

- 1 Myoelectricity produced by intense contractions will interfere with wireless MHR monitoring and computing. It is recommended that you use wired ECG or SpO₂ to monitor MHR during labor.
-

WARNING

- 2 The electrodes for wireless MECG cable should be placed on flat and smooth positions right in the middle of the first intercostals space. Please check the contact of the electrodes regularly.
- 3 When the pregnant woman is lying on the left or right side, please make sure the electrodes are well-placed.

NOTE:

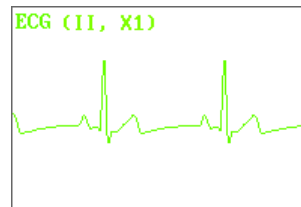
- 1 After the monitor is switched on, if electrodes are not well attached or fell off, alarm message “ECG LEADS OFF” will appear on the screen to draw your attention.
- 2 Vigorous exercise will interfere with MHR monitoring and computing. Please try to avoid it.

9.1.6 Changing ECG Source

Refer to figure 9-2, the ECG signal can come from channel I, II or III. In the ECG trace area of the main interface, **ECG (II, X1)** indicates the ECG source and gain.

If the electrodes are tightly attached to the patient but ECG waveform is not accurate, switch ECG source to another lead by performing the following procedures:

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Lead**.
- 3 Select **I, II** (default) or **III**.
- 4 Select **OK**.



9.1.7 Changing ECG Gain

You can change the ECG gain. The ECG gain affects overall numeric and scope of the ECG waveform.

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Gain**.
- 3 Select **X1/4, X1/2, X1** (default), **X2** or **Auto**.

‘Auto’ means the monitor adjusts the gain automatically. The system displays a 1mv scale at the left side of the ECG waveform. The height of 1mv bar is directly proportional to the waveform amplitude.

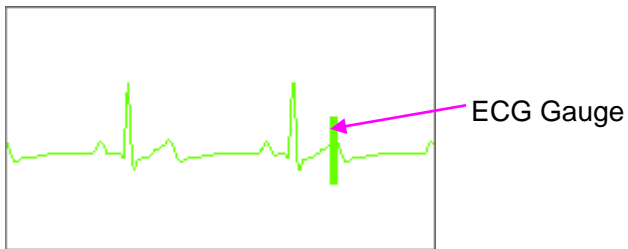
- 4 Select **OK**.

9.1.8 Enabling ECG Calibration

When windage of the ECG waveform is suspected, enable ECG calibration to validate the wave.

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > ECG Calibration**.
- 3 Select **Calibration** or **OFF** (default).
- 4 Select **OK**.

The monitor creates a square wave in the ECG area. Compare the square wave with the ECG gauge. If the error is larger than 0.5mm, change the ECG gain.



When the error is smaller than 0.5mm, calibration is completed. Disable ECG calibration in the same directory.

NOTE:

The ECG Source selection, ECG Gain selection and ECG calibration are only available in the F9 Express monitor with standard configuration.

9.2 Maternal SpO₂ Monitoring

9.2.1 Introduction

The monitor provides continuous monitoring of functional arterial oxygen saturation (SpO₂) and pulse rate (PR) for pregnant women.

SpO₂ Plethysmogram measurement is employed to determine the oxygen saturation of hemoglobin in the arterial blood. If, for example, 97% hemoglobin molecules in the red blood cells of the arterial blood combine with oxygen, then the blood has a SpO₂ oxygen saturation of 97%. The SpO₂ numeric on the monitor will read 97%. The SpO₂ numeric shows the percentage of hemoglobin molecules which have combined with oxygen molecules to form oxyhemoglobin. The SpO₂/PLETH parameter can also provide a pulse rate signal and a plethysmogram wave.

SpO₂ Plethysmogram Measurement Principle:

- ◆ Pulse oximetry is a continuous and noninvasive monitoring technique used to estimate the measurement of arterial oxygen saturation. It measures the amount of light penetrating the patient tissue and reaching the receiver. The reading, obtained through pulse oximetry, uses a light sensor containing two sources of light (red and infrared) that are absorbed by hemoglobin and transmitted through tissues to a photodetector.
- ◆ The amount of light penetrated depends on multiple factors and most of them are constant. However, the arterial blood flow changes with time passing by as is pulsative. The arterial

oxygen saturation can be obtained through testing the absorbed light during pulsation. Plethysmogram wave and pulse rate signal can be also provided during pulsation testing.

The sensor contains LEDs that emit red light at a wavelength of approximately 660 nm and infrared light at a wavelength of approximately 900 nm. Information about wavelength range can be especially useful to clinicians.

The F9 Express monitor is compatible with the SpO₂ transducers supplied by EDAN only. The SpO₂ transducer manufactured by EDAN can only be used with the F9 Express monitor. Compatibility should be checked prior to use. Otherwise the monitor performance can be degraded.

They have been tested and found to comply with the limits for medical device in IEC/EN60601-1-2 (International standard for EMC testing of Medical Electrical Equipment, second edition). These limits are designed to provide reasonable protection against harmful interference in typical medical installation.

WARNING

- 1 Before monitoring, check whether the sensor cable is normal. If any sign of damage in the SpO₂ sensor is detected, do not use the sensor. Return it to the manufacturer for service.
 - 2 Do not put the SpO₂ sensor on the extremities with arterial catheter or venous syringe.
 - 3 Do not perform SpO₂ measuring and NIBP measuring on the same arm at one time, because obstruction of blood flow during NIBP measuring may adversely affect the reading of SpO₂ numeric.
 - 4 Prolonged and continuous monitoring may increase jeopardy of unexpected change of dermal condition such as abnormal sensitivity, rubescence, vesicle, repressive putrescence, and so on. It is especially important to check the sensor placement of neonate and patient of poor perfusion or immature dermogram by light collimation and proper attaching strictly according to changes of the skin.
 - 5 The maximum application time of the SpO₂ sensor at a single site is 4 hours. Check per 2 ~ 3 hours the sensor placement and move it when the skin deteriorates. More frequent examinations may be required for different patients.
-

WARNING

- 6 Setting the SpO₂ higher alarm limit to 100% is equivalent to switching off the alarm on higher limit. High oxygen levels may predispose a premature infant to retrosternal fibroplasia. Therefore, the higher alarm limit for oxygen saturation must be carefully selected in accordance with commonly accepted clinical practices.

CAUTION

Compatibility between the monitor and transducer should be verified before use to avoid injuring the patient or operator.

NOTE:

- 1 The device is calibrated to display functional oxygen saturation.
- 2 A functional tester cannot be used to assess the accuracy of the SpO₂ transducer or the monitor.
- 3 The monitor does not have specific SpO₂ calibration baselines.
- 4 SpO₂ waveform is not proportional to the pulse volume.
- 5 Injected dyes such as methylene blue or intravascular dyshemoglobins such as methemoglobin and carboxyhemoglobin may lead to inaccurate measurements.
- 6 SpO₂ measurement is not applicable during low perfusion and movement.

Measurement Limits -

In operation, the accuracy of oximetry readings can be affected by:

- 1) Magnetic resonance imaging (MRI) scanning. Induced current could potentially cause burns.
- 2) Excessive patient movement.
- 3) Low perfusion.
- 4) High-frequency electrical noise, including noise created by the host system, or noise from external sources, such as electrosurgical apparatus, which is admitted by the host system.
- 5) Intravascular dye injections.
- 6) Improper sensor application.
- 7) Sensor temperature. (Maintain the temperature between +28 °C (+82.4 °F) and +41 °C (+105.8 °F) for best operation)
- 8) Placement of the sensor, such as on an extremity that has a NIBP cuff, arterial catheter, or intravascular line.
- 9) Significant concentrations of dysfunctional hemoglobin, such as carboxyhemoglobin and methemoglobin.

- 10) External illumination more than 5,000 lumens/square meter (typical office lighting). (Cover the sensor site with opaque materials is recommended.)
- 11) Venous pulsations.

To use the sensor:

- a) Select an appropriate sensor. Use the SpO₂ transducers approved by the manufacturer.
- b) Apply the sensor as directed, and observe all warnings and cautions presented in the sensor user manual.
- c) Clean and remove any substances, such as nail polish, from the application site.
- d) Periodically check to ensure that the sensor remains properly positioned on the patient.
- e) Cover the sensor site with opaque material.

9.2.2 SpO₂ Monitoring Procedure

- 1) Insert the SpO₂ sensor plug into the SpO₂ socket on the monitor.
- 2) Place the forefinger, middle finger or third finger into the SpO₂ sensor, refer to figure 9-3.



Figure 9-5 Placement of the Finger

NOTE:

- 1 The nail should cover the light but not too long.
- 2 The cable should be placed on the backside of the hand.
- 3 Avoid external light sources such as radiated rays or ultrared rays.

9.2.3 Enabling SpO₂ Trace Printing

The real-time SpO₂ measurement result is displayed in the parameter area of the main interface. You can choose to print them as a continuous trace on the recorder paper (refer to figure 5-1).

To enable or disable SpO₂ trace printing,

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > SpO₂ Trace**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

9.2.4 Assessing the Validity of a SpO₂ Reading

You can check the quality of the pleth wave and the stability of the SpO₂ values to assess whether the sensor functions properly and whether the SpO₂ readings are valid. Always use these two indications simultaneously to assess the validity of a SpO₂ reading.

NOTE:

- 1 The SpO₂ accuracy has been validated in human studies against arterial blood sample reference measured with a CO-oximeter. Pulse oximeter measurements are statistically distributed, only about two-thirds of the measurements can be expected to fall within the specified accuracy compared to CO-oximeter measurements. The volunteer population in the studies composed of local healthy men and women from age 19 to 37, with variations of skin pigmentations. The SpO₂ accuracy is as follows: $\pm 2\%$ for 90%-100% and $\pm 4\%$ for 70%-90%.
- 2 The pulse rate accuracy is obtained by comparison to the pulse rate generated with an arterial oxygen simulator (also an electronic pulse simulator).
- 3 Generally, the quality of the SpO₂ pleth wave reflects the quality of the light signals obtained by the sensor. A wave of poor quality manifests a decline of the signal validity. On the other hand, the stability of the SpO₂ values also reflects the signal quality. Different from varying SpO₂ readings caused by physiological factors, unstable SpO₂ readings are resulted from the sensor's receiving signals with interference. The problems mentioned above may be caused by patient movement, wrong sensor placement or sensor malfunction. To obtain valid SpO₂ readings, try to limit patient movement, check the placement of the sensor, measure another site or replace the sensor.

9.2.5 SI (Signal Intensity)*

*Only applicable to the EDAN module

The signal intensity (SI) shows perfusion in numeric, and it reflects the pulse intensity of the measurement site. The SI ranges from 0 to 10, with a larger value indicating the more intense signal. When the SI value reaches 10, the signal quality is optimal. If the SI value is less than 2, it indicates that the pulse at the current site is weak, and you should change the measurement site.

The SI value is displayed in the SpO₂ parameter area.



9.2.6 Switching the SpO₂ Alarm On or Off

You can choose to switch the SpO₂ alarm on or off.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **SpO₂**.
- 4 Select **ON** (default) or **OFF** for **Alarm**.
- 5 Select **OK**.

9.2.7 Changing SpO₂ Alarm Limits

You can change the SpO₂ alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **SpO₂**.
- 4 Select a value from 50 ~ 99 for **Low Alarm Limit**. (The step is 1, and the default value is 90%.)
- 5 Select a value from 51 ~ 100 for **High Alarm Limit**. (The step is 1, and the default value is 100%.)
- 6 Select **OK**.

9.3 Maternal HR Monitoring

9.3.1 Introduction

Maternal heart rate (MHR) monitoring does not need an extra accessory. When you perform ECG or SpO₂ (Pulse) monitoring, the MHR result can be acquired at the same time.

When monitoring ECG and SpO₂ at the same time, you can choose the HR source. If only one of them is being performed, the source will automatically switch to the available one (the screen reading should prevail).

9.3.2 Choosing HR Source

You can change the HR source.

- 1 Select the setup key  on the main interface.

- 2 Select **Mother > HR Source**.
- 3 Select **ECG** (default) or **Pulse** (during SpO₂ monitoring).
- 4 Select **OK**.

9.3.3 Changing HR Beep Volume

When the HR beep is enabled, the monitor gives a beep sound of maternal heart. The frequency of HR beep corresponds to the maternal heart rate, but occasionally it may differ due to weak HR signal.

To change the HR beep volume,

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > HR Beep**.
- 3 Select **OFF** (default), **Low** or **High**.
- 4 Select **OK**.

NOTE:

- 1 The DECG beep and HR beep share the same audio channel. Once the HR beep is switched on, the DECG beep is disabled (set to **OFF**) automatically.
- 2 Once the DECG/HR beep volume is changed, the sound switches to channel 1 automatically. Therefore, it is advised against changing DECG/HR beep volume in the monitoring process.

9.3.4 Enabling HR Trace

The real-time MHR measurement result is displayed in the parameter area of the main interface. Also, you can choose to display and print those as a continuous trace on the recorder paper (refer to figure 5-1).

To enable or disable HR trace printing,

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > HR Trace**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

9.3.5 Switching the HR Alarm On or Off

You can choose to switch the HR alarm on or off.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.

- 3 Select **HR**.
- 4 Select **ON** (default) or **OFF** for **Alarm**.
- 5 Select **OK**.

9.3.6 Changing HR Alarm Limits

You can change the HR alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **HR**.
- 4 Select a value from 28 ~ 242 for **Low Alarm Limit**. (The step is 1, and the default value is 50 bpm.)
- 5 Select a value from 29 ~ 243 for **High Alarm Limit**. (The step is 1, and the default value is 120 bpm.)
- 6 Select **OK**.

9.3.7 Signals Overlap Verification

When monitoring maternal heart rate and fetal heart rate at the same time, there are possibilities that maternal HR signal is mistaken for FHR signal. The SOV function of the monitor can also reduce these possibilities.

In the process of monitoring, if the SOV detects signals overlapping, an alarm message “Signals Overlap (FHR1/FHR2/DFHR, HR)” will appear on the screen to warn you. Checking the patient and reposition of transducers might be needed.

9.4 Maternal NIBP Monitoring

9.4.1 Introduction

The monitor measures blood pressure using the oscillometric method.

Oscillometric devices measure the amplitude of pressure changes in the occluding cuff as the cuff deflates from above systolic pressure. The amplitude suddenly increases as the pulse breaks through the occlusion in the artery. As the cuff pressure decreases further, the pulsations increase in amplitude, reach a maximum (which approximates to the mean artery pressure), and then diminish.

There are two modes available: Manual and Auto. In manual mode, NIBP is measured once on each demand. In auto mode, NIBP is measured repeatedly after a preset time interval. This interval is adjustable. You can perform a manual measurement during an Auto measurement interval.

In both modes, systolic pressure (SYS), diastolic pressure (DIA) , mean artery pressure (MAP)

and pulse rate (PR) (optional) are measured and displayed.

The blood pressure measurements determined with this device comply with the American National Standard for Electronic or Automated Sphygmomanometers (ANSI/AAMI/ISO 81060-2:2013) in relation to mean error and standard deviation.

In clinical investigation method with a reference sphygmomanometer, the fifth Korotkoff sound was used to determine adult diastolic pressure, and the fourth Korotkoff sound was used to determine pediatric diastolic pressure.

WARNING

- 1 Check for any fault of the cuff before start monitoring.
- 2 Do not perform NIBP measurements on patients with sickle-cell disease or under any condition where the skin is damaged or expected to be damaged, such as on the arm on the side of a mastectomy.
- 3 Pressurization of the cuff can temporarily cause loss of function of simultaneously used monitor on the same limb.
- 4 If liquid is splashed on or into the main unit inadvertently, or enters the conduit, stop using the monitor and contact the manufacturer for service immediately.
- 5 For a thrombasthenia patient, it is important to determine whether the measurement of blood pressure shall be done automatically. The determination should be based on clinical evaluation.
- 6 Do not apply the cuff to a limb that has an intravenous infusion or catheter in place frequently. This could cause tissue damage around the catheter when infusion is slowed or blocked during cuff inflation.
- 7 Make sure that the air conduit connecting the blood pressure cuff and the monitor is neither blocked nor tangled.

NOTE:

The monitor is intended to measure NIBP for adults only.

Measurement Limitations -

To different patient conditions, the oscillometric measurement has certain limitations. The measurement is in search of regular arterial pressure pulse. In those circumstances where the patient's condition makes it difficult to detect, the measurement becomes unreliable and the measuring time increases. You should be aware that the following conditions could interfere with the measurement, making the measurement unreliable or longer to derive. In some cases, the patient's condition will make a measurement impossible.

1) Patient Movement

Measurements will be unreliable or may not be possible if the patient is moving, shivering or having convulsions. These motions may interfere with the detection of the arterial pressure pulses. In addition, the measurement time will be prolonged.

2) Cardiac Arrhythmia

Measurements will be unreliable and may not be possible if the patient's cardiac arrhythmia has caused an irregular heartbeat. The measuring time thus will be prolonged.

3) Heart-lung Machine

Measurements will not be possible if the patient is connected to a heart-lung machine.

4) Pressure Changes

Measurements will be unreliable and may not be possible if the patient's blood pressure is changing rapidly over the period of time during which the arterial pressure pulses are being analyzed to obtain the measurement.

5) Severe Shock

If the patient is in severe shock or hypothermia, measurements will be unreliable since reduced blood flow to the peripheries will cause reduced pulsation of the arteries.

6) Heart Rate Extremes

Measurements can not be done to a patient whose heart rate is lower than 40 bpm or higher than 240 bpm.

9.4.2 How to Apply NIBP Cuff

WARNING

Accuracy of NIBP measurement depends on using a cuff of the proper size. It is essential to measure the circumference of the limb and choose the proper cuff size. If you find something is wrong with the cuff size, please replace it immediately.

- 1) Select appropriate cuff for the patient.

The table below lists the reference size:

Type	Limb Perimeter	Cuff Size	Air Hose Length
Upper Arm (Adult 1)	27 cm ~ 35 cm	14.5 cm	3 m
Upper Arm (Adult 2)	34 cm ~ 43 cm	18 cm	

- 2) Squeeze the cuff to discharge the air.

- 3) Apply the cuff to the patient; make sure that the index line is placed in the appointed range and

↑
the symbol ↑ is over the appropriate artery (Refer to figure 9-4). If the index line is not in the appointed range, please replace for a proper one. Ensure that the cuff is not wrapped too tightly around the limb. Excessive tightness may cause discoloration and eventual ischemia of the extremities.



Figure 9-6: Applying the Cuff

9.4.3 Preparation for NIBP Monitoring

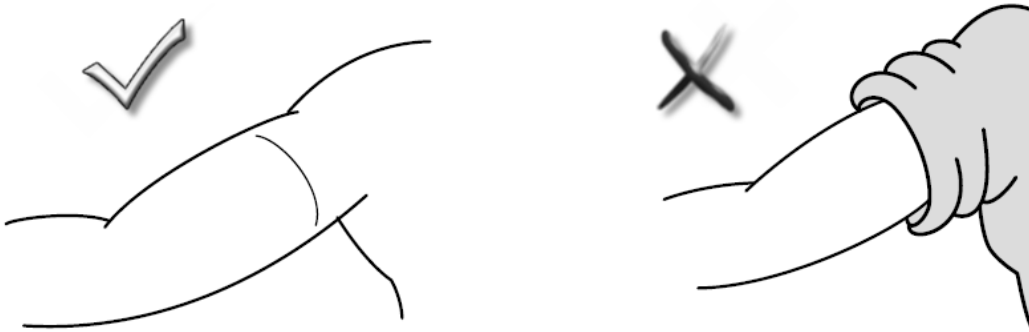
To obtain accurate measurements, the following operating steps need to be observed:

1. Ensure the patient position in normal use, including
 - ◆ comfortably seated
 - ◆ legs uncrossed
 - ◆ feet flat on the floor
 - ◆ back and arm supported
 - ◆ middle of the cuff at the level of the right atrium of the heart
2. Relax as much as possible and do not talk during the measurement.
3. Wait for five minutes until the first reading is taken.

NOTE:

Please roll up the sleeve and keep the patient's arm bare or it will cause the inaccurate measurements.

- 1) Wrap the cuff on a bare arm.



- 2) Insert the cuff plug into NIBP socket on the monitor.
- 3) Apply the NIBP cuff to the patient's arm or leg following the instructions described in section 9.4.2 How to Apply NIBP Cuff.

- 4) Connect the cuff to the air hose. The limb chosen for taking the measurement should be placed at the same level as the patient's heart. If this is not possible, correct the measurement using the formula described in section 9.4.6 *Correcting the Measurement*.



Figure 9-7 Connection for NIBP measurement

9.4.4 *Auto Measurement

To perform an auto measurement,

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Cycle**.
- 3 Select a time interval from 1, 2, 3, 4, 5, 10, 15, 30, 60, 90, 120, 180, 240 and 480 minutes.
- 4 Select **OK**.
- 5 Press **NIBP** key on the front panel to start an Auto measurement.

NOTE:

The monitor checks uterine contract (UC) pressure when the **NIBP** key is pressed. If the UC is over 50, a prompt “Intense UC, can't measure NIBP now.” is issued, and the monitor will check the UC every 30 seconds. The monitor will measure NIBP only when the UC is lower than 50, and it will then start timing for the Auto measurement.

To stop the current measurement,

Press the **NIBP** key anytime during the current measurement to stop it. Another measurement will start after the time interval.

WARNING

Prolonged NIBP measurements in automatic mode may be associated with purplish patches, ischemia and neurologic damage in the limb wearing the cuff. When monitoring a patient, examine the extremities of the limb frequently for normal color, warmth and sensitivity. If any abnormality is observed, stop the NIBP measurement.

9.4.5 *Manual Measurement

To perform a manual measurement,

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Cycle**.
- 3 Select **Manual**.
- 4 Select **OK**.
- 5 Press **NIBP** key on the front panel to start a manual measurement.

To stop the manual measurement,

Press the **NIBP** key anytime during the measurement to stop it.

To perform a manual measurement during an auto measurement interval,

1 Press the **NIBP** key to start the manual measurement.

2 Press the **NIBP** key again anytime to stop it.

The monitor will restart timing for the Auto measurement and resume measuring after the time interval.

NOTE:

- 1 If you are in doubt about the accuracy of any reading(s), check the patient's vital signs by an alternative method before checking the functioning of the monitor.
- 2 The monitor checks uterine contract (UC) pressure when the **NIBP** key is pressed. If the UC is over 50, a prompt "Intense UC, can't measure NIBP now." is issued. Please wait and do not attempt to measure NIBP until the UC is lower than 50.

CAUTION

- 1 Do not squeeze the rubber tube on the cuff.
- 2 If liquid is inadvertently splashed on the equipment or its accessories, or may enter the conduit or inside the monitor, contact local service center.

9.4.6 Correcting the Measurement

To correct the measurement if the limb is not at heart level,

- ◆ add 0.75 mmHg (0.10 kPa) for each inch higher.
- ◆ deduct 0.75 mmHg (0.10 kPa) for each inch lower.

9.4.7 Changing NIBP Unit

You can change the NIBP unit.

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Unit (NIBP Setup)**.
- 3 Select **mmHg** (default) or **kPa**.

- 4 Select **OK**.

Note:

When the network version is ETHERNET 1.4, if the fetal monitor gets on line, the central station will control to make the NIBP unit of the fetal monitor in concert with that of the central station.

9.4.8 Switching the NIBP Alarm On or Off

You can choose to switch the NIBP alarm on or off. The SYS alarm and DIA alarm are related. Once one of them is switched off, the rest will be switched off as well.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **SYS, DIA or MAP**.
- 4 Select **ON** (default) or **OFF** for **Alarm**.
- 5 Select **OK**.

9.4.9 Changing SYS Alarm Limits

You can change the SYS alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **SYS**.
- 4 Select a value from 40 ~ 269 (mmHg) or 5.3~35.9 (kPa) for **Low Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 90 mmHg; if the unit is kPa, the step is 0.1, and the default value is 12.0 kPa.)
- 5 Select a value from 41 ~ 270 (mmHg) or 5.4~36.0 (kPa) for **High Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 160 mmHg; if the unit is kPa, the step is 0.1, and the default value is 21.3 kPa.)
- 6 Select **OK**.

9.4.10 Changing DIA Alarm Limits

You can change the DIA alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **DIA**.
- 4 Select a value from 10 ~ 214 (mmHg) or 1.3~28.6 (kPa) for **Low Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 50 mmHg; if the unit is kPa, the step is 0.1, and the default value is 6.8 kPa.)

- 5 Select a value from 11 ~ 215 (mmHg) or 1.4~28.7(kPa) for **High Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 90 mmHg; if the unit is kPa, the step is 0.1, and the default value is 12.0 kPa.)
- 6 Select **OK**.

9.4.11 Changing MAP Alarm Limits

You can change the MAP alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **MAP**.
- 4 Select a value from 20 ~ 234 (mmHg) or 2.7 ~31.2 (kPa) for **Low Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 60 mmHg; if the unit is kPa, the step is 0.1, and the default value is 8.0 kPa.)
- 5 Select a value from 21 ~ 235 (mmHg) or 2.8~31.3 (kPa) for **High Alarm Limit**. (If the unit is mmHg, the step is 1, and the default value is 110 mmHg; if the unit is kPa, the step is 0.1, and the default value is 14.8 kPa.)
- 6 Select **OK**.

9.4.12 *Choosing NIBP Printing Mode

When the recorder is printing real-time fetal traces, the NIBP result is also recorded on the paper whenever NIBP measurement is performed. After the paper stops advancing, you can choose to keep recording NIBP results on the paper.

To enable or disable NIBP printing after paper advancing stops,

- 1 Select the setup key  on the main interface.
- 2 Select **Recorder > NIBP**.
- 3 Select **ON** or **OFF** (default).
- 4 Select **OK**.

NOTE:

- 1 The NIBP measurement results during the period of paper lacking and fast printing after new paper is loaded will not be printed. Therefore, do not perform NIBP measurements during this period.
- 2 NIBP measurement can be affected by extremes of temperature, humidity and altitude.

9.4.13 *Calibrating NIBP

NIBP is not user-calibrated. Cuff-pressure transducers must be verified and calibrated, if necessary, at least once every two years by a qualified service professional.

9.5 Maternal TEMP Monitoring

9.5.1 TEMP Monitoring Procedure

- 1) Insert the TEMP plug into the TEMP socket on the monitor.
- 2) Apply the sensor firmly underneath the patient's axilla. It takes 5 minutes for the temperature measurement to stabilize.

WARNING

- 1 Check if the TEMP sensor functions properly prior to use.
- 2 Do not apply the TEMP sensor to the mouth or the rectum.

CAUTION

Be cautious when taking and putting the TEMP sensor. Do not pull the cable too tight or it might cause mechanical damage.

The transient response time for the continuous TEMP sensor is not larger than 30s. The laboratory method used to test the response time is as follows:

1. Prepare two reference temperature sources. The first one with a constant temperature of 25 °C (77 °F) and the second one with a constant temperature of 27 °C (80.6 °F).
2. Put the TEMP sensor to the first reference temperature source until the temperature reading reaches 25 °C (77 °F).
3. Move the TEMP sensor to the second reference temperature source. Note the time (t1) from the TEMP sensor being moved in to the temperature reading reaching 27 °C (80.6 °F).
4. When the temperature reading is stable, move the TEMP sensor back to the first reference temperature source. Note the time (t2) from the TEMP sensor being moved in to the temperature reading falling to 25 °C (77 °F).
5. The larger value of t1 and t2 is the response time.

Note :

The reference body site temperature is the same as the temperature of the measuring site.

9.5.2 Changing TEMP Unit

You can change the TEMP unit.

- 1 Select the setup key  on the main interface.
- 2 Select **Mother > Unit (TEMP Setup)**.

- 3 Select **°C** (default) or **°F**.
- 4 Select **OK**.

Note:

When the network version is ETHERNET 1.4, if the fetal monitor gets on line, the central station will control to make the TEMP unit of the fetal monitor in concert with that of the central station.

9.5.3 Switching the TEMP Alarm On or Off

You can choose to switch the TEMP alarm on or off.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **TEMP**.
- 4 Select **ON** (default) or **OFF** for **Alarm**.
- 5 Select **OK**.

9.5.4 Changing TEMP Alarm Limits

You can change the TEMP alarm limits.

- 1 Select the setup key  on the main interface.
- 2 Select **Alarm**. On the displayed **Password** box, enter **9999**, then select **Enter**.
- 3 Select **TEMP**.
- 4 Select a value from 0.0 ~ 49.9 for **Low Alarm Limit**. (The step is 0.1, and the default value is 36.0 °C.)
- 5 Select a value from 0.1 ~ 50.0 for **High Alarm Limit**. (The step is 0.1, and the default value is 39.0 °C.)
- 6 Select **OK**.

Chapter 10 Maternal Monitoring Display (F9 Express)

10.1 *Display Mode

F9 Express has three display modes: maternal-fetal display (figure 10-1), fetal display (figure 10-2) and maternal display (figure 10-3).

To change the display mode, select the display mode switch  on the main interface. The display mode will switch among the three modes.

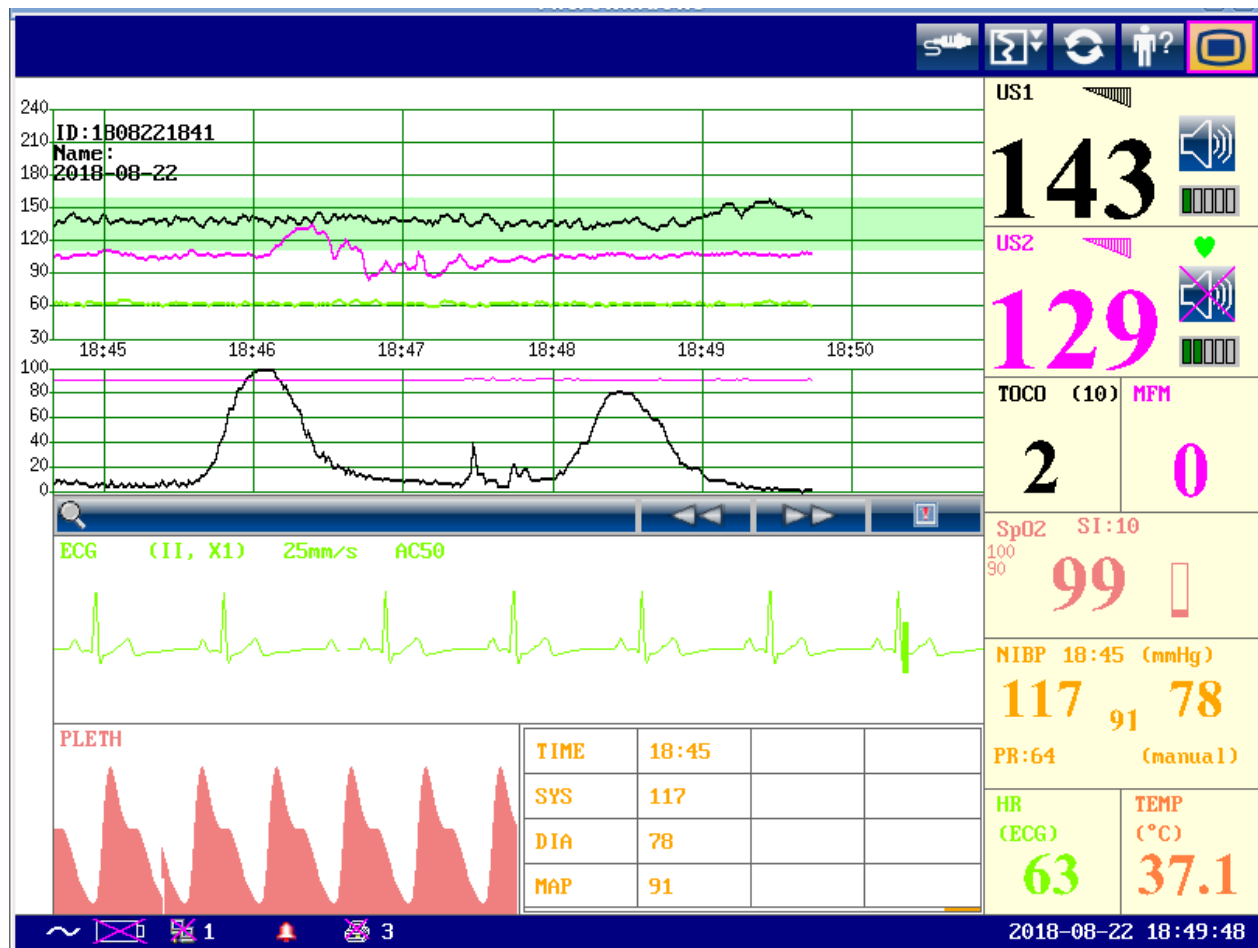


Figure 10-1 Maternal-Fetal Display mode

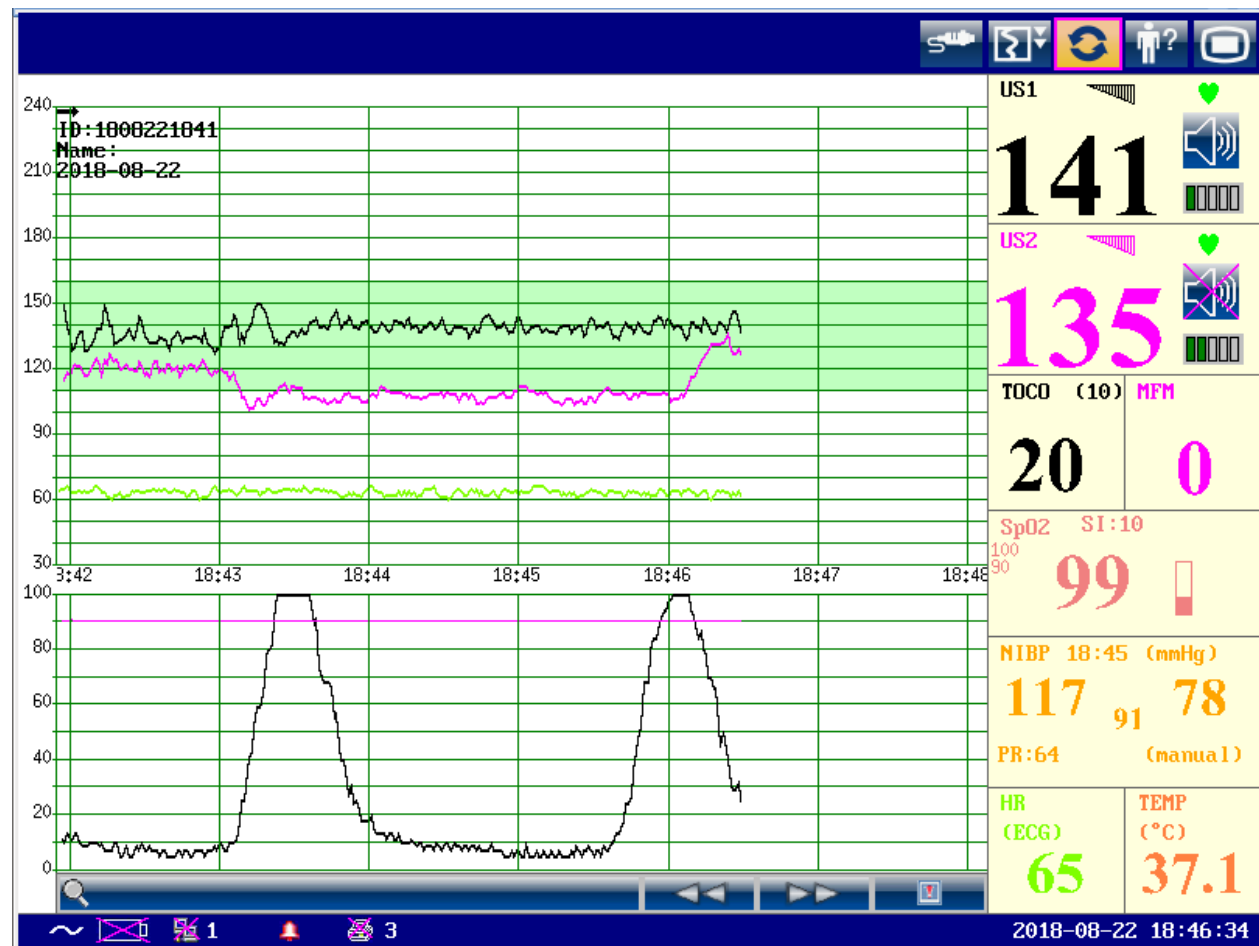


Figure 10-2 Fetal Display Mode

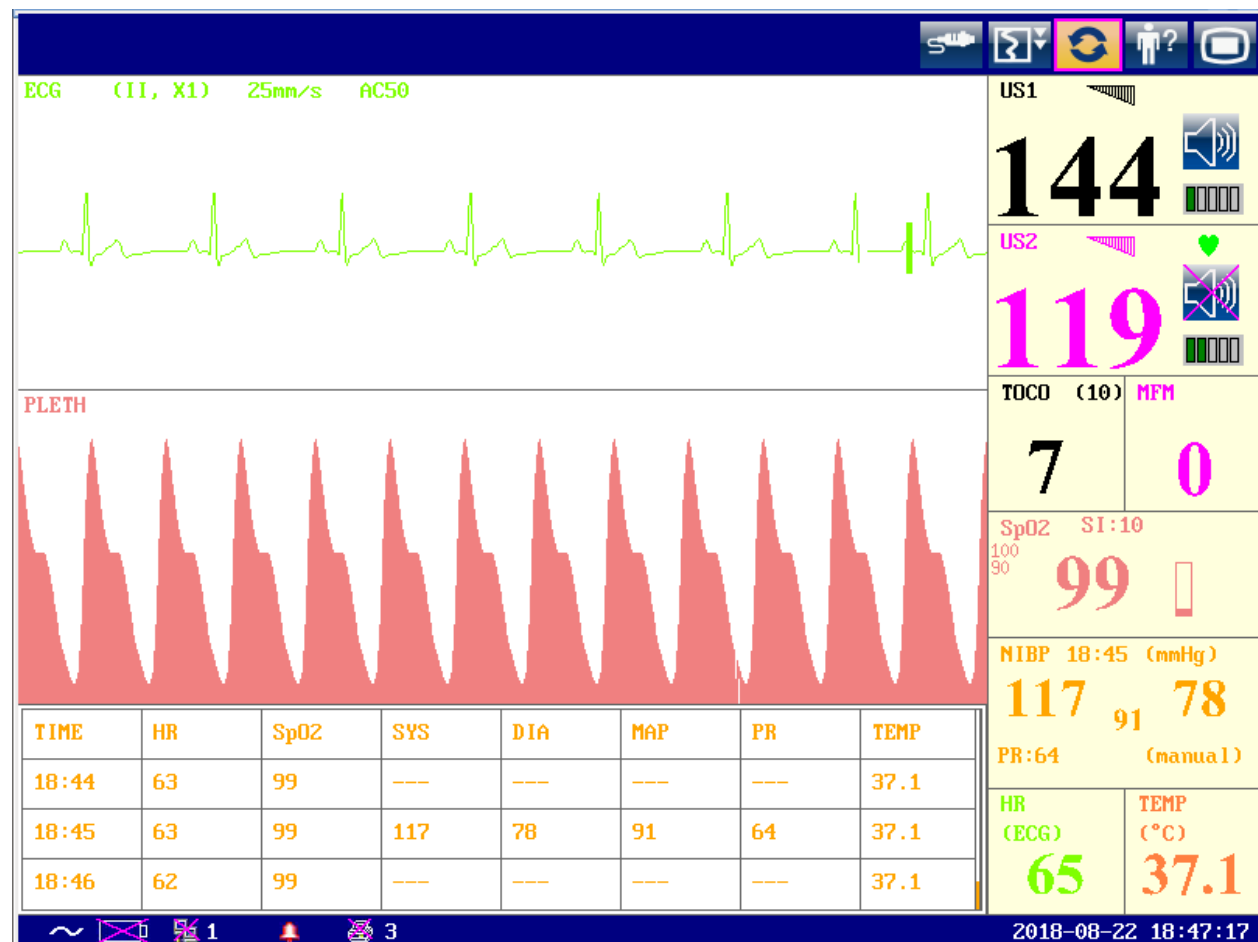


Figure 10-3 Maternal Display Mode

10.2 Maternal Monitoring Traces

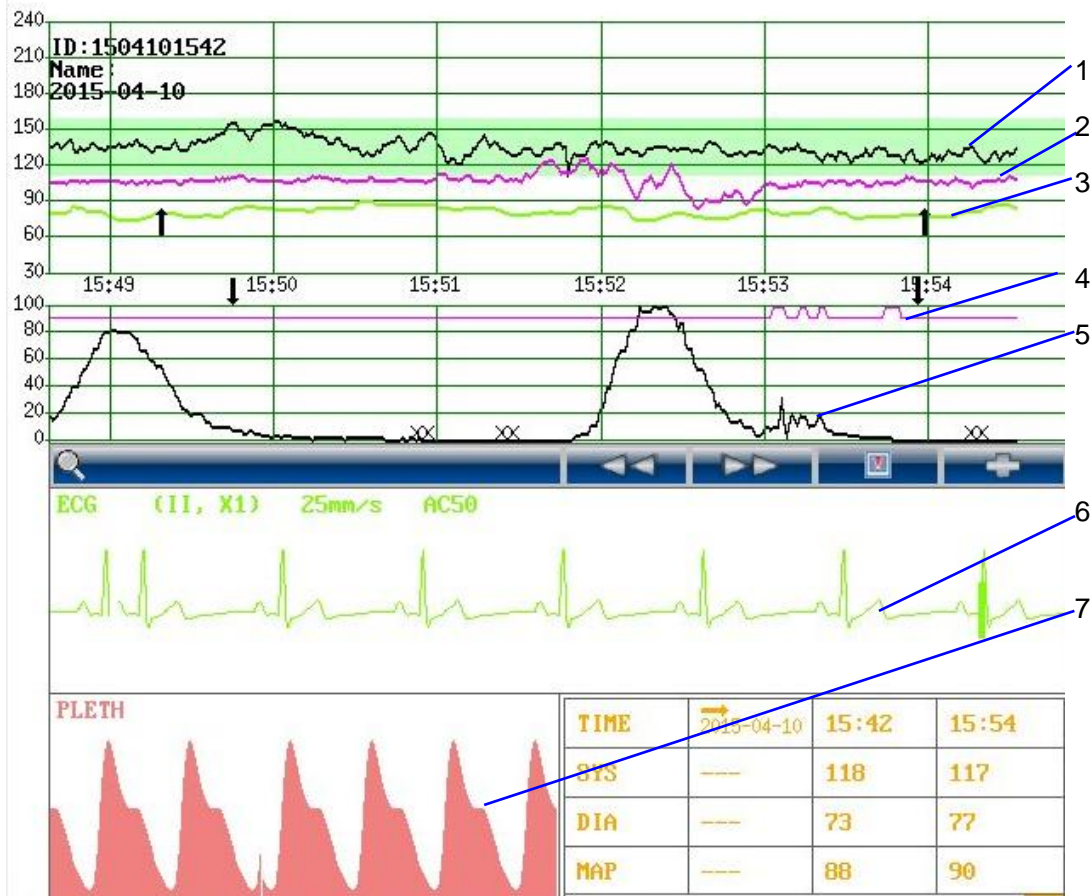


Figure 10-4 Maternal Monitoring Traces

- | | | | |
|--------------|--------------|-----------------------------|-------------|
| 1 FHR1 Trace | 2 FHR2 Trace | 3 HR Trace | 4 AFM Trace |
| 5 TOCO Trace | 6 ECG Trace | 7 SpO ₂ Waveform | |

F9 Express displays both maternal monitoring traces and fetal monitoring traces on the same screen. The maternal monitoring traces include ECG waveform and SpO₂ waveform. The fetal monitoring traces are the same as traces of **F9**, refer to *8.1 Traces* for more information.

10.3 Maternal Vital Sign List

The maternal vital sign list keeps records of the recent maternal vital signs and the measuring time. A start mark → and the date appear when a new monitoring begins.

In maternal-fetal display mode, the list contains the time, SYS, DIA and MAP numerics of every measurement.

TIME	2015-03-30	14:40	14:42
SYS	---	118	117
DIA	---	73	77
MAP	---	88	90

Figure 10-5 Maternal NIBP List

In maternal display mode, the list contains the time, HR, SpO₂, SYS, DIA, MAP, PR and TEMP numerics. The numerics are recorded every minute.

TIME	HR	SpO2	SYS	DIA	MAP	PR	TEMP
16:56	69	99	114	78	90	60	37.1
16:57	68	99	---	---	---	---	37.1

Figure 10-6 Maternal Vital Sign List

The numeric list can be reviewed: select the list, press the symbol or rotate the control knob to review the previous lists.

10.4 Numerics

Besides the fetal numerics, the numeric window of **F9 Express** includes maternal vital signs: SpO₂, NIBP, HR and TEMP:

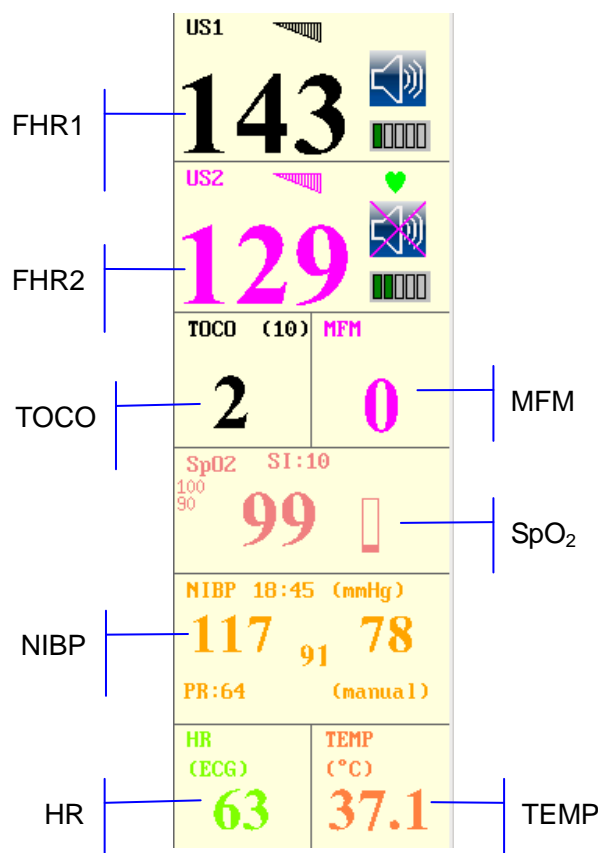


Figure 10-7 Maternal Numeric Window

SpO₂ 	99: Current SpO ₂ measurement numeric.
	100: SpO ₂ alarm upper limit
	90: SpO ₂ alarm lower limit
	■: SpO ₂ indicator. SI: Signal intensity.
NIBP 	18:45: Time when the NIBP measurement starts.
	mmHg: NIBP unit.
	From left to right, top to bottom in turn: current systolic pressure (117), mean artery pressure (91), diastolic pressure (78) and PR (64).
	(manual): The current NIBP measurement mode is manual.
HR 	(ECG): The current HR comes from ECG.
	63: Current maternal heart rate measurement numeric.

TEMP	(°C): TEMP unit.
TEMP 37.1	

When F9 Express Fetal/Maternal Monitor is connected to FTS-3 Telemetry System, the signal strength and signal quality icon and battery level icon of the US-T transducers and TOCO-T transducer or TOCO-E transducer are displayed in the numeric window.

If wireless AFM and wireless MECG are enabled, the signal strength icon is also displayed for them.

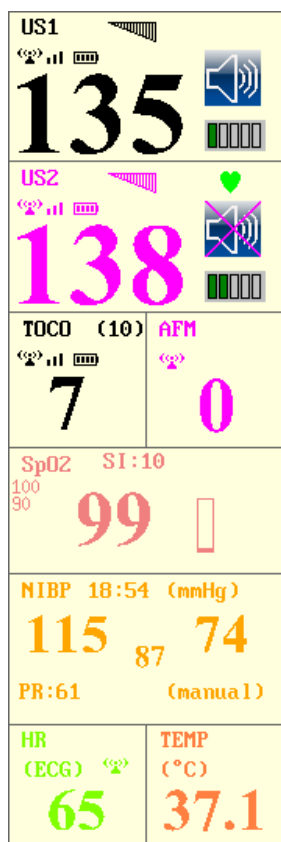


Figure 10-8

If FHR1 is monitored using wireless DECG (a TOCO-E transducer connected with DECG cable), the signal strength and signal quality icon and battery level icon displayed in the DECG numeric window are the TOCO-E transducer's signal strength and battery level. Please note that wireless MECG cannot be used while wireless DECG is in use.

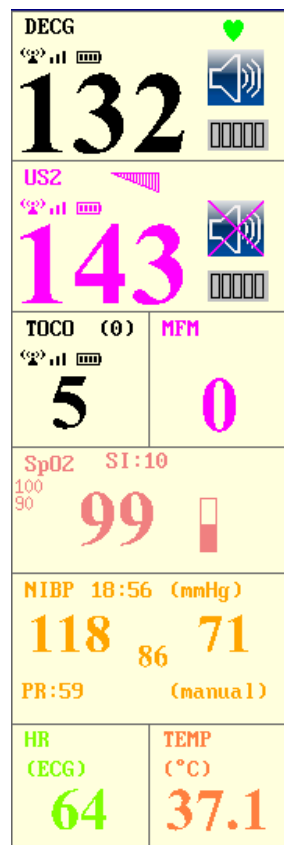


Figure 10-9

Chapter 11 After Monitoring

11.1 Completing Monitoring

After monitoring,

- 1) Remove transducers or electrodes from the patient; wipe the remaining gel off the patient and the transducer with a clean soft cloth or tissue.
- 2) Press the **PRINT** key to stop printing, and press the paper advancing key  to advance the paper.
- 3) Wait the paper to stop and then tear it off along the perforation.

NOTE:

After the fetus is delivered in the labor, the monitor may pick up signals of the umbilical cord and display a trace/numeric. To avoid misinterpretation, it is recommended to remove the transducers from the patient and switch off the monitor immediately after the fetus is delivered.

11.2 Switching Off

- 1) Press and hold the **POWER** switch for at least 3 seconds to switch off the monitor.
- 2) Unplug the power cord.

CAUTION

Do not press the POWER switch continuously. Allow at least 10 seconds between switching the monitor on and off.

Chapter 12 Maintenance and Cleaning

12.1 Maintenance

12.1.1 Maintaining Inspection

(1) Visual Inspection

Prior to using the monitor or FTS-3 every time, do the following inspections:

- ◆ Check the monitor and accessories to see if there is any visible evidence of damage that may affect patient safety. Pay special attention to the cracks on the transducers and cables before immersing them into conductive fluid.
- ◆ Check all the outer cables, power socket and power cables.
- ◆ Check if the monitor functions properly.

If any damage is detected, stop using the monitor or FTS-3 on the patient. Replace the damage part(s) or contact the manufacturer for service before reusing it.

(2) Routine Inspection

The overall check of the monitor and the accessories, including safety check and function check, should be performed by qualified personnel every 6 to 12 months, and each time after service.

The equipment should undergo periodic safety testing to ensure proper patient isolation from leakage currents. This should include leakage current measurement and insulation testing. The recommended testing interval is once a year or as specified in the institution's test and inspection protocol.

(3) Mechanical Inspection

Make sure all exposed screws are tight.

Check the external cables for splits, cracks or signs of twisting.

Replace any cable that shows serious damage.

Pay particular attention to the supply socket.

WARNING

Failure on the part of the responsible individual hospital or institution employing the use of this equipment to implement a satisfactory maintenance schedule may cause undue equipment failure and possible health hazards.

CAUTION

Besides the maintenance requirements recommended in this manual, comply with local regulations on maintenance and measurement.

12.1.2 Maintenance of Monitor and Base Station

Keep the exterior surface of the monitor and the base station clean, free of dust and dirt.

The gathering of dew on the screen may occur with abrupt temperature or humidity changes. A stable environment is recommended. Stop using the monitor or the base station and contact the service personnel immediately if accidental wetting occurs.

Scratching and damaging the screen should be avoided.

Operate the touch screen with special stylus pen or finger. Sharp edged or hard particles like ball pen or propelling pencil are prohibited. Keep the touch screen surface clean, and no adhesive should be applied. Avoid high voltage and static charge.

12.1.3 Maintenance of Wired and Wireless Transducers

WARNING

- 1 The transducers must be cleaned before docking in the base station after each use. Make sure that there is no residual coupling gel.
- 2 The transducers are delicate and sensitive. Please handle them with care and try to avoid dropping on to the ground or any hard surfaces.

Although transducers are designed for durability, they should be handled with care. Rough handling could damage the cover, piezoelectric crystals and mechanical movement. Contacting the transducers with hard or sharp objects should be avoided. Do not excessively flex the cables.

The transducers must be thoroughly cleaned and disinfected at least once a month. When cleaning, please firstly use a lint-free cloth moistened with mild near neutral detergent, ethanol 75% solution or isopropanol 70% alcohol-based solution to clean the transducers. Then use a cotton cloth moistened with clear water to clean again. At last, use a dry, soft cloth to dry them.

In case of unsuccessful charge or poor contact, please use detergent with abrasive effect to rub the electrodes of the transducers in order to clear away the oxide of coupling gel.

Charge and discharge the wireless transducer battery every 3 months.

12.1.4 Storage of Recorder Paper

When storing recorder paper (including used paper with traces):

Do not store in plastic envelopes.

Do not leave exposed to direct sunlight or ultraviolet light.

Storage conditions outside these limits may distort the paper and adversely affect the accuracy of grid lines or make the trace unreadable.

12.1.5 Cleaning of Recorder

The recorder platen, thermal print head and paper sensing mechanism must be cleaned at least once a year or when needed (when traces become faint).

To do this:

- 1) Clean the recorder platen with a lint-free cloth dampened in soap/ water solution.
- 2) Wipe the thermal array using a cotton swab moistened with 70% Isopropyl alcohol-based solution.
- 3) Check that the paper sensing mechanism is free of dust.

WARNING

Switch off the monitor and remove the power cord prior to recorder cleaning.

12.1.6 Maintaining the Battery

It is required to follow the instructions in this user manual during installation, storage and maintenance of the battery.

When the battery is charged, used or stored, keep it away from objects or materials with static electric charges.

The recommended charge temperature range is from 0 °C (+32 °F) to +40 °C (+104 °F). Do not exceed this range.

When not using battery for a long time, remove it from the monitor and store it in a place with low humidity and low temperature.

Batteries have life cycles. If the time that the monitor uses the battery becomes much shorter than usual, the battery life is at an end. Replace it with a new one the same as the one provided or recommended by the manufacturer.

12.2 Cleaning

In order to avoid infection, clean and disinfect the monitor and accessories after each use.

12.2.1 Cleaning of Monitor and Base Station

Regular cleaning of the monitor enclosure and the screen is strongly recommended.

WARNING

- 1 Unplug the monitor and the base station from the AC power source and detach all accessories before cleaning. Do not immerse the unit in water or allow liquids to enter the case.
- 2 If liquid is splashed on or into the main unit inadvertently, or enters the conduit, stop using the monitor and contact the manufacturer for service immediately.

The solutions recommended for monitor cleaning are: mild near neutral detergent, ethanol 75% and isopropanol 70%.

Clean the monitor and the base station enclosure with soft cloth and diluent non-caustic detergents recommended above.

Clean the screen and the charging point in the docking slot with a dry soft cloth.

CAUTION

- 1 Although the monitor and the base station are chemically resistant to most common hospital cleaners and non-caustic detergents, different cleaners are not recommended and may stain the monitor.
- 2 Many cleansers must be diluted before use. Follow the manufacturer's directions carefully to avoid damaging the monitor and the base station.
- 3 Do not use strong solvent, for example, acetone.
- 4 Never use an abrasive such as steel wool or metal polish.
- 5 Do not allow any liquid to enter the product, and do not immerse any part of the monitor into any liquid.
- 6 Avoid pouring liquids on the monitor while cleaning.
- 7 Do not allow any remaining solution on the surface of the monitor.

NOTE:

- 1 The monitor surface can be cleaned with hospital-grade ethanol and dried in air or with crisp and clean cloth.
- 2 The manufacturer has no responsibility for the effectiveness of controlling infectious disease using these chemical agents. Please contact infectious disease experts in your hospital for details.

12.2.2 Cleaning of Accessories

(1) Cleaning of Transducers

To clean the transducers and leads, follow these steps:

- 1) Wipe them with a soft cloth dampened in cleaning solution;
- 2) Clean them with a soft cloth dampened in water;

3) Air-dry them or wipe the remaining moisture with a soft dry cloth.

The recommended cleansers for accessories are listed below:

Accessory	Cleansers
Ultrasound Transducer TOCO Transducer (Including the wireless)	Mild near neutral detergent Ethanol 75% Isopropanol 70%
DECG Leads	Mild near neutral detergent Ethanol 75% Isopropanol 70%
IUP Cable	Mild near neutral detergent Ethanol 75% Isopropanol 70%
ECG Leads	Mild near neutral detergent Ethanol 75% Isopropanol 70%
SpO ₂ Transducer	Mild near neutral detergent Ethanol 75% Isopropanol 70%
TEMP Transducer	Mild near neutral detergent Ethanol 75% Isopropanol 70%

CAUTION

- 1 The waterproof parts of the transducer are restricted to the main body and the cable.
Do not immerse the plug into water during the process of monitoring or cleaning.
- 2 Be sure the temperature of cleaning solutions does not exceed +45 °C (+113 °F).
- 3 Only wipe the outer surface of accessories. Do not immerse them in any liquid.
- 4 Make sure no liquid enters the connector.
- 5 When you clean the TEMP transducer, take the head in one hand and clean with the soft cloth in the other hand.
- 6 After cleaning, no remaining cleanser is allowed on the surface.
- 7 Please clean the charging point periodically or it will not be charged.

(2) Cleaning of Belt

Wash soiled belts with soap and water. The water temperature must not exceed +60 °C (+140 °F).

(3) Cleaning of NIBP Cuff

The cuff can also be machine-washed or hand-washed. Hand-washing will prolong the life of the cuff.

Remove the latex rubber bag before washing; for machine-washing, close the Velcro fastening. Allow the cuff to dry thoroughly after washing; then reinsert the rubber bag.

Replace the Rubber Bag in the Cuff

To replace the rubber bag in the cuff, first place the bag on the top of the cuff so that the rubber tubes line up with the large opening on the long side of the cuff. Now roll the bag lengthwise and insert it into the opening on the long side of the cuff. Hold the tubes and the cuff and shake the complete cuff until the bag is in position. Thread the rubber tubes from inside the cuff, and out through the small hole under the internal flap.

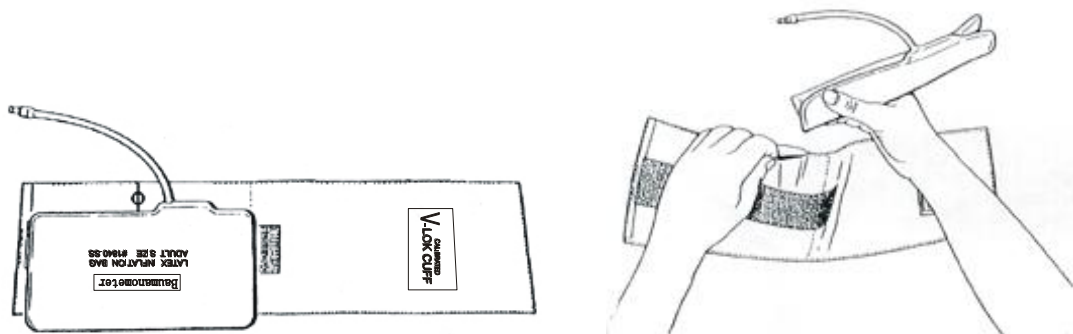


Figure 12-1 Replace the Rubber Bag in the Cuff

CAUTION

- 1 Do not squeeze the rubber tube on the cuff.
- 2 Do not dry-clean the cuff.
- 3 Only clean the outer surface of the connectors, make sure no liquid goes into the connector.
- 4 When the reusable cuff is not connected with the monitor, or being cleaned, always place the cover on the rubber tube to avoid liquid permeation.

12.3 Disinfecting

To disinfect the transducers and leads, follow these steps:

- 1) Clean the accessories.
- 2) Wipe them with a soft cloth dampened in the recommended disinfectant.
- 3) Wipe them clean with a soft cloth dampened in water.
- 4) Air-dry them or wipe the remaining moisture with a soft dry cloth.

The table below lists the allowed disinfectant bases:

Type	Recommended
Fetal/Maternal Monitor	
Base Station	
US and TOCO Transducers (the wired and wireless)	Ethanol 75%
Remote Event Marker	Isopropanol 70%
DEC G Cable	
IUP Cable	
ECG Leads	
SpO ₂ Transducer	
TEMP Transducer	

Type	Recommended
NIBP Cuff	Ethanol 75%
NIBP Cuff Extension Tube	Isopropanol 70%

CAUTION

- 1 Do not use any disinfectant containing additional active ingredients other than those listed.
- 2 Follow the manufacturer's instruction to dilute the solution, or adopt the lowest possible density.
- 3 Do not immerse any part of the monitor or any accessory into liquid.
- 4 After disinfection, no remaining disinfectant is allowed on the surface.
- 5 Check if the monitor and accessories are in good condition. If any aging or damage is detected (e.g. the belt loses its elasticity), replace the damage part(s) or contact the manufacturer for service before reusing them.
- 6 Please do not light the TOCO transducer with ultraviolet light for a long time.

NOTE:

The manufacturer has no responsibility for the effectiveness of controlling infectious disease using these chemical agents. Please contact infectious disease experts in your hospital for details.

12.4 Sterilizing

Do not sterilize the monitor, the base station or the accessories, unless this is necessary according to your hospital regulation.

NOTE:

Check if the monitor, base station, cables and accessories function well. If any problem is detected, please contact the manufacturer for service before reusing them.

Checking Item	Checking Method
Visual	Inspect the monitor, base station and cables etc. for any damage.
Power On	Power on the monitor. Does it boot up successfully without errors and enter the main menu?
Functionality Test	After power up, check whether the AC power indicator and battery status indicator in the bottom left of the screen display as stated in section 2.4.1.
Performance	Please check the US transducer and TOCO transducer according to section 7.2.6 Testing US Transducers and section 7.5.4 Testing TOCO Transducers. FTS-3 wireless transducers also can be tested accordingly.
System	When the monitor is connected to FTS-3, please check whether the base station working channel and its battery status indicator in the bottom right of the screen display as stated in section 2.4.1.

Chapter 13 Warranty and Service

13.1 Warranty

EDAN warrants that EDAN's products meet the labeled specifications of the products and will be free from defects in materials and workmanship that occur within warranty period.

The warranty is void in cases of:

- a) damage caused by mishandling during shipping.
- b) subsequent damage caused by improper use or maintenance.
- c) damage caused by alteration or repair by anyone not authorized by EDAN.
- d) damage caused by accidents.
- e) replacement or removal of serial number label and manufacture label.

If a product covered by this warranty is determined to be defective because of defective materials, components, or workmanship, and the warranty claim is made within the warranty period, EDAN will, at its discretion, repair or replace the defective part(s) free of charge. EDAN will not provide a substitute product for use when the defective product is being repaired.

13.2 Contact information

If you have any question about maintenance, technical specifications or malfunctions of devices, contact your local distributor.

Alternatively, you can send an email to EDAN service department at: support@edan.com.cn.

Appendix 1 Product Specifications

A F9, F9 Express Fetal/Maternal Monitor

A1.1 Environmental Specifications

The monitor may not meet the performance specifications given here if stored or used outside the specified temperature and humidity ranges.

Working	Temperature	+5 °C ~ +40 °C (+41 °F ~ +104 °F)
	Relative Humidity	15% ~ 93% (non-condensing)
	Atmospheric Pressure	86 kPa ~ 106 kPa
Transport and Storage	Temperature	-20 °C ~ +55 °C (-4 °F ~ +131 °F)
	Relative Humidity	15% ~ 93% (non-condensing)
	Atmospheric Pressure	70 kPa ~ 106 kPa

A1.2 Physical Specifications

Monitor		
Dimensions and Weight	Size (depth x width x height)	347mm x 330mm x 126mm
	Weight	F9: Approx. 5.5 kg F9 Express: Approx. 6.3 kg
Power Supply	Operating Voltage	100V-240V~
	Operating Frequency	50Hz/60Hz
	Input Power	1.0A-0.5A
	Battery	14.8VDC/5000mAh
Standards Compliance	IEC 60601:2005+A1:2012, EN 60601-1:2006+A1:2013, IEC 60601-1-2:2014, EN 60601-1-2:2015, IEC/EN 60601-2-27, IEC/EN 60601-2-37, IEC/EN 60601-2-49, IEC 80601-2-30, ISO 80601-2-61, ISO 80601-2-56, AAMI/ANSI EC13	
Anti-electric Shock Type	Class I equipment with internal power supply	

Anti-electric Shock Degree	FHR1, FHR2, TOCO, FM, IUP SpO ₂ , NIBP DECG ECG, TEMP	BF BF (Defibrillating-proof) CF CF (Defibrillating-proof)
Degree of Protection against Harmful Ingress of Water	Main Unit: IPX1, protected against vertically falling water drops (provided recorder drawer is shut and the monitor is not mounted on the wall vertically) US/TOCO Transducers: IPX8, protected against the effects of continuous emersion in water	
Degree of Safety in Presence of Flammable Gases	Equipment not suitable for use in presence of flammable gases	
Disinfection/Sterilizing Method	Refer to this user manual for details	
EMC	CISPR11 Group 1 Class A	
Working System	Continuous running equipment	
Display		
Screen Diagonal	12.1"	
Pixel	800(H) × 600(V)	
Signal Interface		
RS232 interface (DB9 or D-Sub), RJ45 interface		
Ultrasound Transducer		
Cable Length	2.5m	
Weight	190 g	
Dimension	88 mm × 35 mm	
TOCO Transducer		
Cable Length	2.5 m	
Weight	180 g	
Dimension	88 mm × 35 mm	
Remote Event Marker		
Length	2.5 m	
Weight	56 g	
ECG		

Cable Length	3 m
Weight	213 g
SpO₂	
Cable Length	2.4 m
Weight	68 g
NIBP	
Cable Length	3.3 m
Weight	194 g
TEMP	
Cable Length	3 m
Weight	55 g

A1.3 Performance Specifications

FHR	*FHR Measurement Range	50 bpm ~ 240 bpm
	*Resolution	1 bpm
	*Accuracy	±1 bpm
	*Alarm	FHR Alarm
	*Ultrasound Output	Isppa.3<190W/cm ²
		Ispta.3<94mW/cm ²
		Isata<20 mW/cm ²
		TI<1.0 MI<1.0
	*Temperature Rise	When applied to the patient, the ultrasound transducer may warm slightly (less than 2 °C (3.6 °F) above ambient temperature). When NOT applied, at the ambient temperature of 40 °C (104 °F), the ultrasound transducer may reach the highest temperature of 43 °C (109.4 °F).
	p- <1 MPa	
	I _{ob} <10 mW/cm ²	
	I _{spta} <100 mW/cm ²	
	Max Output Power <15mW	
	Effective Radiating Area	(942 ± 15%) mm ²
	Dielectric Strength	4000Vrms
TOCO	*TOCO Range	0~100

	*Non-linear Error	$\pm 10\%$
	*Resolution	1
	Baseline Drift due to Temperature Changes	1 unit/min/ °C (free air) 5 units/min/ °C (underwater)
	Zero Mode	Automatic (TOCO value becomes zero or below lasting for 30 seconds)/ Manual
	Dielectric Strength	4000Vrms
DECG	*DFHR Measurement Range	30bpm ~ 240bpm
	*Resolution	1bpm
	*Accuracy	$\pm 1\text{bpm}$
	*Alarm	DFHR Alarm
	Technique	Peak-peak detection technique
	Input Impedance	> 10M (Differential, DC50/60Hz)
	Input Impedance	> 20M (Common Mode)
	CMRR	> 110dB
	Noise	< 4 μVp
	Skin Voltage Tolerance	$\pm 500\text{mV}$
IUP	Fetal Input Voltage Current	20 μVp -3mVp
	*Pressure Range	0mmHg ~100mmHg (0.0 kPa~13.3 kPa)
	*Non-linear Error	$\pm 3\text{mmHg}$ ($\pm 0.4 \text{ kPa}$)
	*Resolution	1mmHg (0.1 kPa)
	Sensitivity	5 $\mu\text{V/V/mmHg}$
MFM AFM &	Zero Mode	Manual
	*Display Range	0 ~ 999
	*FM Mode	Automatic/Manual
	*AFM Mode	Trace (default) or Black Mark
MECG	AMF Technique	Pulsed Doppler ultrasound
	*MHR Measurement Range	30bpm ~ 240bpm

*MHR Measuring Accuracy	$\pm 2\text{bpm}$
*Resolution	1 bpm
* MHR Alarm Limits	30bpm ~ 240bpm
*Alarm	HR Alarm
*Anti-electric Shock Type	Defibrillating-proof
Input Signal Range	$\pm 8 \text{ mV PP}$
ECG Waveform	Manual control ECG waveform display
ECG falls off	Detect automatically
Patient Leakage Current (Limit)	N.C. S.F.C. d.c. $10 \mu\text{A}$ $50 \mu\text{A}$ a.c. $10 \mu\text{A}$ $50 \mu\text{A}$
Patient Auxiliary Current (Limit)	N.C. S.F.C. d.c. $10 \mu\text{A}$ $50 \mu\text{A}$ a.c. $10 \mu\text{A}$ $50 \mu\text{A}$
Differential Input Impedance	$>5\text{M}\Omega$
Display Sensitivity	2.5mm/mV ($\times 0.25$), 5mm/mV ($\times 0.5$), 10mm/mV ($\times 1$), 20mm/mV ($\times 2$), AUTO gain
Electrode Offset Potential Tolerance	$\pm 500\text{mV}$
Auxiliary Current (Leads off detection)	Active electrode: $< 100 \text{nA}$ Reference electrode: $< 900 \text{nA}$
Accuracy and Response to Irregular Rhythm	According with ANSI/AAMI EC13-2002 Sect.4.1.2.1 e) The MHR value displays after a stable period of 20s: Ventricular bigeminy: $80\text{bpm} \pm 1\text{bpm}$ Slow alternating ventricular bigeminy: $60\text{bpm} \pm 1\text{bpm}$ Rapid alternating ventricular bigeminy: $120\text{bpm} \pm 1\text{bpm}$ Bidirectional systoles: $91\text{bpm} \pm 1\text{bpm}$
Bandwidth(-3dB)	Diagnosis: 0.05 Hz to 150 Hz Monitor: 0.5 Hz to 40 Hz
Response time to Change in MHR	MHR range: 80bpm ~ 120bpm

		Range: 7s ~ 8s (average: 7.5s) MHR range: 80bpm ~ 40bpm Range : 7s ~ 8s (average: 7.5s)
	Tall T-wave Rejection	Exceeds ANSI/AAMI EC13-2002 Sect. 3.1.2.1 (C) minimum recommended 1.2mV T-Wave amplitude
SpO2	*Measurement Range	50% ~ 100%
	*Resolution	1 %
	*Measuring Accuracy (EDAN)	90% ~ 100% ±2% 70% ~ 90% ±4% < 70% unspecified
	*Measuring Accuracy (Nellcor)	70% ~ 100% ±2% < 70% unspecified
	*Data update period (EDAN)	1s
	*Data update period (Nellcor)	2s
	*PR Measurement	Range: 30~240bpm
		Resolution: 1 bpm
		Accuracy: ±3 bpm
	*SpO2 Alarm Limits	50% ~ 100%
	*Alarm	PR Alarm and SpO2 Alarm
	Wavelength	Red light: (660±3) nm
		Infrared light: (905±10) nm
		Emitted light energy: < 15 mW
	Information about the wave length range can be especially useful to clinicians (for instance, when photodynamic therapy is performed.)	
NIBP	*Measurement	Systolic Pressure, Diastolic Pressure, Mean Artery Pressure
	*Method	Oscillometric Method
	*Measurement Range	Systolic Pressure: 40 mmHg ~ 270 mmHg (5.3 kPa~36.0 kPa) Diastolic Pressure: 10 mmHg ~ 215 mmHg (1.3

		kPa~28.7 kPa) Mean Artery Pressure: 20 mmHg ~ 235 mmHg (2.7 kPa~31.3 kPa)
*Resolution	1 mmHg (0.1 kPa)	
*Measuring Accuracy	Max. average deviation $\leq \pm 5\text{mmHg}$ ($\leq \pm 0.8\text{kPa}$) Max. standard deviation $\leq 8\text{mmHg}$ ($\leq 1.2\text{kPa}$)	
*Measuring Time (Normal)	30 ~ 45s	
*Measuring Time (MAX)	120s	
*Alarm Limits	Systolic Pressure: 40 mmHg ~ 270 mmHg (5.3 kPa~36.0 kPa) Diastolic Pressure: 10 mmHg ~ 215 mmHg (1.3 kPa~28.7 kPa) Mean Artery Pressure: 20 mmHg ~ 235 mmHg (2.7 kPa~31.3 kPa)	
*Alarm	Systolic Pressure, Diastolic Pressure, Mean Artery Pressure Alarm	
Software Over Voltage Protection	(297 \pm 3) mmHg [(39.6 \pm 0.4) kPa]	
Hardware Over Voltage Protection	(320 \pm 10) mmHg [(42.8 \pm 1.3) kPa]	
Cuff pressure measuring range	0 mmHg ~ 300 mmHg (0.0 kPa~40.0 kPa)	
TEMP	*Channel	1
	*Measurement Range	0°C ~ +50°C
	*Resolution	0.1°C
	*Accuracy	$\pm 0.3\text{ }^{\circ}\text{C}$ (Transducer error excluded: $\pm 0.1\text{ }^{\circ}\text{C}$) (Transducer: $\leq \pm 0.2\text{ }^{\circ}\text{C}$)
	*Unit	°C, °F
	*Refresh Time	1 ~ 2s
	*Self Check	5 ~ 10min
	*Alarm Limits	0.0°C ~ +50.0°C
	*Alarm	TEMP Alarm
	Measuring Mode	Direct Mode

	Position	Axilla
	Accessory	TEMP transducer

NOTE:

The essential performance is marked with an asterisk *.

A1.4 Recorder Specifications

Paper	Z-fold, thermosensitive (compatible with GE and PHILIPS recorder papers)
Paper width	152mm (GE), 150mm (PHILIPS)
Effective printing width	110mm (American Standard) 120mm (International Standard)
FHR printout width	70mm (American Standard) 80mm (International Standard)
FHR scaling	30bpm/cm (American Standard) 20bpm/cm (International Standard)
TOCO printout width	40mm
TOCO scaling	25%/cm
Printing speed	
Standard Speed(Real-Time Traces)	1 cm/min, 2 cm/min, 3 cm/min
Fast Print Speed(Stored Traces)	Up to 15mm/sec
Accuracy of data	±5% (X axis)
Accuracy of data	±1% (Y axis)
Resolution	8 dots/mm
Record Information	FHR1 trace/mark, FHR2 trace/mark, TOCO trace, AFM trace/black mark, fetal movement mark, event mark (and annotation), AUTO-zero symbol, alarm indicator, SOV alarm indicator, US1 and US2 signal loss alarm indicator, wired/wireless monitoring status mark, date, time, printing speed, ID, name, FHR2 Offset, HR, SpO ₂ , SYS, DIA, MAP, PR, TEMP, CTG analysis results etc.

A1.5 Rechargeable Lithium-ion Battery

Type	Rechargeable Lithium-ion Battery
Continual Working Time	>2 hours
Necessary charge time from "out of power" to "fully charged"	<7hours
Necessary charge time from "out of power" to "90% charged"	<5 hours
Nominal Capacity	5000mAh
Nominal Voltage	14.8V
Cycle Life	> 300 times

A1.6 Low Output Summary Table

Low Output Summary Table

(for systems with no transducers having global maximum index values exceeding 1.0)

System: Fetal & Maternal Monitor

Transducer Model	I _{spta.3} (mW/cm ²)	TI Type	TI Value	MI	I _{pa.3@MI_{max}} (W/cm ²)
PW1.0MHz	1.92	TIS	0.0091	0.013	0.011
		TIB	0.051		

B FTS-3 Fetal Telemetry System

B1.1 Environmental Specifications

Working	Temperature	+5 °C ~ +40 °C (+41 °F ~ +104 °F)
	Relative Humidity	15% ~ 93% (non-condensing)
	Atmospheric Pressure	86 kPa ~ 106 kPa
Transport and Storage	Temperature	-20 °C ~ +55 °C (-4 °F ~ +131 °F)
	Relative Humidity	15% ~ 93% (non-condensing)
	Atmospheric Pressure	70 kPa ~ 106 kPa

B1.2 Physical Specifications

Power Supply	Operating Voltage	100V-240V~
	Operating Frequency	50Hz/60Hz
	Input Power	0.8A-0.3A
	Battery	14.8VDC/5000mAh
Standards Compliance	IEC 60601:2005+A1:2012, EN 60601-1:2006+A1:2013, IEC 60601-1-2:2014, EN 60601-1-2:2015, IEC/EN 60601-2-37, IEC 60601-2-27, EN 62479:2010, ETSI EN 301 489-1, ETSI EN 301 489-3, ETSI EN 300 220-1, ETSI EN 300 220-2.	
Anti-electric Shock Type	Class I equipment with internal power supply	
Anti-electric Shock Degree	FHR1, FHR2, TOCO DECG, MHR(from MECG)	BF CF
Degree of Protection against Harmful Ingress of Water	Base station: IPX1 (protected against vertically falling water drops) Transducers: IPX8 (Protected against the effects of continuous immersion in water 1.1m deep for 24 hours)	
Degree of Safety in Presence of Flammable Gases	Equipment not suitable for use in presence of flammable gases	
Disinfection/Sterilizing Method	Refer to this user manual for details	
EMC	CISPR11 Group 1 Class A	

Leakage Current			
Ground Leakage Current (Limit)	N.C. 500 µA	S.F.C. 1000 µA	
Enclosure Leakage Current (Limit)	N.C. 100 µA	S.F.C. 500 µA	
Patient Leakage Current (Limit)	N.C. d.c. 10 µA	S.F.C. 50 µA	
FHR1, FHR2, TOCO	a.c. 100 µA	500 µA	
Patient Auxiliary Current (Limit)	N.C. d.c. 10 µA	S.F.C. 50 µA	
FHR1, FHR2, TOCO	a.c. 100 µA	500 µA	
Patient Leakage Current (Limit)	N.C. d.c. 10 µA	S.F.C. 50 µA	
DECG, MHR(from MECG)	a.c. 10 µA	50 µA	
Patient Auxiliary Current (Limit)	N.C. d.c. 10 µA	S.F.C. 50 µA	
DECG, MHR(from MECG)	a.c. 10 µA	50 µA	
Base Station			
Weight	About 1.8 kg		
Size	310mm x 235mm x 81mm		
US-T Transducer			
Weight	About 150 g		
Size	Ø81 mm x 35 mm		
TOCO-T Transducer & TOCO-E Transducer			
Weight	About 150 g		
Size	Ø81 mm x 35 mm		

B1.3 Performance Specifications

Ultrasound	
*FHR Measurement Range	50 bpm ~ 240 bpm
*Resolution	1 bpm
*Accuracy	± 2 bpm
Technique	Ultrasound Pulse Doppler with autocorrelation
Pulse Repetition Rate	2 kHz
Pulse Duration	92 μ s
Ultrasound Frequency	(1 \pm 10%) MHz
$p_{\perp} < 1 \text{ MPa}$	
$I_{ob} < 10 \text{ mW/cm}^2$	
$I_{spta} < 100 \text{ mW/cm}^2$	
Dielectric Strength	4000 Vrms
TOCO	
*TOCO Range	0~ 100
*Non-linear Error	$\pm 10\%$
*Resolution	1
Baseline Drift due to Temperature Changes	1 unit/min/ $^{\circ}\text{C}$ (free air) 5 units/min/ $^{\circ}\text{C}$ (underwater)
Zero Mode	Automatic/ Manual
Dielectric Strength	4000 Vrms
RF Index	
Transmission Power	Wireless transducer:< 1mW e.r.p Base station:< 10mW e.r.p
Frequency Range	433.050MHz~434.790MHz

*Transmission Range (Without Obstacles)	>110m (when performing underwater monitoring using US-T and TOCO-T transducers, keep the transducer at a distance ≤30cm from the water surface and a distance ≤8m from the base station.)	
Modem Mode	GFSK	
Transmission Rate	About 25kbps	
Channel Range	1~14	
Transducer Antenna	FM antenna	
Base Station Antenna	Internal antenna	
DECG	Technique	Peak-peak detection technique
	*DFHR Measurement Range	30bpm ~ 240bpm
	*Resolution	1bpm
	*Accuracy	±1bpm
	Input Impedance	> 10MΩ (Differential, DC50/60Hz)
	Input Impedance	> 20MΩ (Common Mode)
	CMRR	> 110dB
	Skin Voltage Tolerance	±500mV
	Fetal Input Voltage Current	20 μV-6mV
MHR	* MHR Measurement Range	30 bpm ~ 240 bpm
	Input Signal Range	±8 mV PP
	* MHR Measuring Accuracy	±2 bpm
	* MHR Alarm Limits	30 bpm ~ 240 bpm
	*Anti-electric Shock Type	Defibrillating-proof
	ECG falls off	Detect automatically
	Patient Leakage Current (Limit)	N.C. S.F.C. d.c. 10 μA 50 μA a.c. 10 μA 50 μA

	Patient Current (Limit)	N.C. d.c. 10 µA a.c. 10 µA	S.F.C. 50 µA 50 µA
	Differential Input Impedance	>5MΩ	
	Electrode Offset Potential Tolerance	±500mV	
	Auxiliary Current (Leads off detection)	Active electrode: < 100 nA Reference electrode: < 900 nA	
	Accuracy and Response to Irregular Rhythm	According with ANSI/AAMI EC13-2002 Sect.4.1.2.1 e) The MHR value displays after a stable period of 20s: Ventricular bigeminy: 40bpm±2 bpm Slow alternating ventricular bigeminy: 30~62bpm Rapid alternating ventricular bigeminy: 60bpm±1bpm Bidirectional systoles: 60~100bpm	
	Response time to Change in MHR	MHR range: 80bpm ~ 120bpm Range: 6s ~ 8s (average: 7.2s) MHR range: 80bpm ~ 40bpm Range : 7s ~ 9s (average: 7.6s)	
	Tall T-wave Rejection	Exceeds ANSI/AAMI EC13-2002 Sect. 3.1.2.1 (C) minimum recommended 1.2mV T-Wave amplitude	
AFM	*Display Range	0 ~ 999	
	*FM Mode	Automatic	
	*AFM Mode	Trace (default) or Black Mark	
	Technique	Pulsed Doppler ultrasound	

NOTE:

The essential performance is marked with an asterisk *.

B1.4 Rechargeable Lithium-ion Battery

Base Station Battery	
Nominal Capacity	5000 mAh
Continuous Work Time	≥40 Hours
Nominal Voltage	14.8 V
Necessary Charge Time	≤14 Hours
Cycle Life	>300 times

Transducer Battery	
Nominal Capacity	1600 mAh
Charge Current (Standard)	700 mA
Continuous Work Time	>17h (full new battery used in transducer) >12h (full new battery used in TOCO-E transducer connected with DECG or MECG cable)
Nominal Voltage	3.7 V
Charge Voltage (Standard)	(4.2±0.1) V
Cycle Life	≥500 times

B1.5 Low Output Summary Table

Low Output Summary Table

(for systems with no transducers having global maximum index values exceeding 1.0)

System: Fetal Telemetry System

Transducer: 12-Crystal Wafer

Transducer Model	I _{pta.3} (mW/cm ²)	TI Type	TI Value	MI	I _{pa.3@MI_{max}} (W/cm ²)
PW1.0MHz	1.66	TIS	0.0079	0.017	0.0092
		TIB	0.064		

Low Output Summary Table

(for systems with no transducers having global maximum index values exceeding 1.0)

System: Fetal Telemetry System

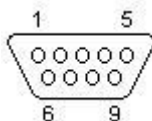
Transducer: 7-Crystal Wafer

Transducer Model	$I_{spta.3}$ (mW/cm ²)	TI Type	TI Value	MI	$I_{pa.3@MI_{max}}$ (W/cm ²)
PW1.0MHz	3.17	TIS	0.050	0.030	0.020
		TIB	0.020		

Appendix 2 Signal Input/ Output Connector

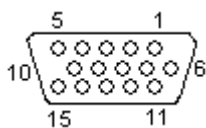
Accessory equipment connected to these interfaces must be certified according to the respective IEC/EN standards (e.g. IEC/EN 60950 for data processing equipment and IEC/EN 60601-1 for medical equipment). Furthermore all configurations shall comply with the valid version of the system standard IEC/EN 60601-1. Anybody who connects additional equipment to the signal input connector or signal output connector to configure a medical system must ensure that the system complies with the requirements of the valid version of the system standard IEC/EN 60601-1. If in doubt, contact our technical service department or your local distributor.

DB9 Interface

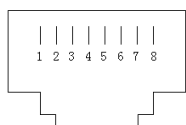


Pin	Signal	Input/Output
1	+5V	Output
2	Rx	Input
3	Tx	Output
4	485EN	Input
5	0V Ref.	
6	TA	Output
7	TB	Output
8	RA	Input
9	RB	Input

D-Sub Interface



Pin	Signal	Input/Output
1	US2	Input
2	ISOCNS_RXD	Input
3	ISOCNS_TXD	Output
4	485EN	Input
5	0V Ref.	
6	TA	Output
7	TB	Output
8	RA	Input
9	RB	Input
10	DECG_SIGNAL	Input
11	US1	Input
12	+5V	Output
13	TOCO	Input
14	DECG_GND	
15	EN	Input

RJ45 Interface

Pin	Signal	Input/Output
1	TD+	Output
2	TD-	Output
3	RD+	Input
4	Reserved	
5	Reserved	
6	RD-	Input
7	Reserved	
8	Reserved	

CAUTION

Only the PC or telemetry system recommended by the manufacturer can be connected to the signal input/output interface of the monitor. Other equipment is forbidden.

Appendix 3 Troubleshooting

A F9, F9 Express Fetal/Maternal Monitor

A3.1 No Display

Phenomenon	Possible Cause	Solution
Power indicator is off.	Power cable is loose.	Tighten the power cable.
	The fuse is blown.	Change the fuse.
	The battery runs out of power.	Connect to AC power supply.

A3.2 Noise

Phenomenon	Possible Cause	Solution
Noise	Too high volume.	Turn down the volume.
	Interfered by mobile phone or other interfering source.	Keep the interfering source far away from the monitor.

A3.3 Recorder Error

Phenomenon	Possible Cause	Solution
Paper jam	Wrong loading paper or paper is dampened.	Load paper correctly and keep paper from moist.
Recorder does not work.	The recorder is not started.	Press the PRINT key.
	Run out of paper.	Load paper.
	The paper drawer is not locked.	Slide the paper drawer in until both latches are locked in position.
Incorrect time and date	Time incorrectly set	Reset time and date and note the difference between Daylight Saving Time and Winter Standard Time(See 6.5)
	Battery fault	The battery needs service. Call the service personnel.

A3.4 Trouble with Ultrasound FHR Monitoring

Phenomenon	Possible Cause	Solution
Inconstant trace / display	The patient is overweighted.	Monitor FHR with DECG.
	Improper ultrasound transducer position.	Adjust the position of the transducer till the better signal is received.
	Loose belt.	Tighten the belt.
	Superfluous aquasonic coupling gel.	Wipe off superfluous aquasonic coupling gel.
	Frequent fetal movements.	Delay the monitoring.
	Maternal movement.	Request the patient to calm down and stay still.
	Inadequate aquasonic coupling gel.	Use recommended aquasonic coupling gel quantity.
Doubtful FHR	Record maternal heart rate wrongly.	Change the position of the ultrasound transducer.
	The transducer is not well placed in position, and the mixed noise has been recorded.	Adjust the position of the transducer.
Faint trace or no trace	Improper paper.	Use paper recommended by manufacturer
	The paper drawer is not locked.	Slide the paper drawer in until both latches are locked in position.
	Adjusting nuts of the print head are unbalanced.	Contact the manufacturer for service.

A3.5 Troubles with DECG Monitoring

Phenomenon	Possible Cause	Solution
Inconstant trend Inconstant display	No ECG signal	Use a new spiral electrode
	Bad contact of reference electrode and patient	Use a new spiral electrode
Inconstant trend	The DECG cable has not been fixed firmly	Fix an attachment pad at the DECG cable.

A3.6 Troubles with Contractions Monitoring (External)

Phenomenon	Possible Cause	Solution
Bad trace quality or fluctuant TOCO baseline	The belt is too tight or too loose.	Adjust the belt.
	The belt has no elasticity.	Renew the belt.
	Maternal movement.	Request the patient to calm down and stay still.
	Frequent fetal movements.	Delay the monitoring.
Too high TOCO sensitivity (higher than 100 unit)	The body pressure from uterus to TOCO transducer is far higher than the average numeric.	Insure favorable contact for patient skin with TOCO transducer. Change the position of TOCO transducer, if necessary.

A3.7 Troubles with Contractions Monitoring (Internal)

Phenomenon	Possible Cause	Solution
No trend	The intrauterine catheter is jammed	Wash with disinfectant
No pressure change when uterine contraction	“Dry” environment or the tip of intrauterine catheter is placed extraovularly	Wash with disinfectant or change the position of transducer
Only see the IUP peak but no baseline	Zero adjustment is wrong	Zero the system
The trend is a beeline	The connector failure.	Move or contact catheter. If trend no fluctuation, change intrauterine cable.

A3.8 Big ECG Signal Interference or Thick Baseline

Phenomenon	Possible Cause	Solution
Big ECG signal interference or thick baseline	Abnormal electrodes placing or electrodes invalidation.	Check the electrodes placing and the period of validity of electrodes.
	The cable connector is not well connected.	Check the connection of cable connector.
	Power socket has no standard ground wire.	Check if power socket has standard ground wire.

	The special ground wire connecting with monitor is not properly earthed.	Check if the special ground wire connecting with monitor is earthed.
--	--	--

A3.9 NIBP and SpO2 No Results

Phenomenon	Possible Cause	Solution
NIBP and SpO ₂ have no results	The NIBP cuff is not properly wrapped to the position of patient's arm.	Check if the NIBP cuff is properly wrapped to the position of patient's arm.
	The NIBP can not be inflated.	Extend catheter, and check the connection.
	Hose connector plug is not connected well with the NIBP socket.	Check if the hose connector plug is connected well with the NIBP socket.
	SpO ₂ transducer is not connected well with the SpO ₂ socket.	Check if the SpO ₂ transducer is connected well with the SpO ₂ socket.
	Abnormal working condition.	Shut off the power, then switch it on again.

A3.10 Blown Fuses

WARNING

Switch off the monitor and unplug it before changing the fuse.

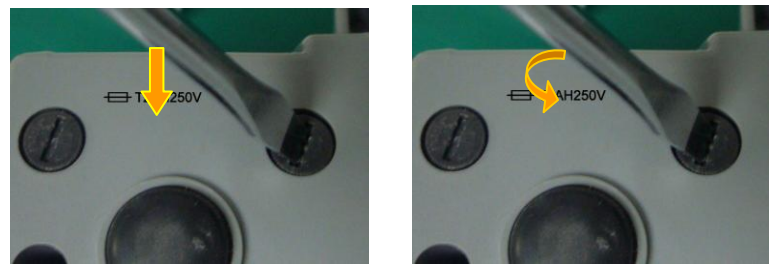
Replace the fuse when it is blown.

The two fuses of the monitor are located on the bottom panel, their specifications are:

Size: Φ5mm*20mm; Model: T2AH250V.

To replace a fuse:

- 1) Fold the LCD display completely flat.
- 2) Carefully place the monitor upside down on a flat surface covered with cloth or other protecting pad.
- 3) With a flat-head screw driver, push the fuse in for about 1 mm and then unscrew it anticlockwise.
- 4) Remove the old fuse and replace it with a new fuse that is supplied by the manufacturer or of the same specifications.
- 5) Push the new fuse into the socket for about 1 mm and then screw it clockwise back in position.



B FTS-3 Fetal Telemetry System

B3.1 Troubleshooting

Phenomenon	Possible Cause	Solution
Take out the US-T transducer, but it cannot power on.	<ul style="list-style-type: none"> ① It runs out of power. ② The base station cannot communicate with the transducer by RF. 	<ul style="list-style-type: none"> ① Recharge the transducer. ② Put it back in the docking slot and take it up again. If the problem persists, restart the base station.
The wireless connection indicator is green but the fetal monitor shows no signal.	<ul style="list-style-type: none"> ① Loose or damaged cable to the monitor socket 	<ul style="list-style-type: none"> ① Tighten or repair the cable.
FHR or TOCO record interrupts.	<ul style="list-style-type: none"> ① Transducer is placed incorrectly. ② Transducer slides. ③ The patient walks in strong tramps. ④ RF interference or out of prescriptive area. 	<ul style="list-style-type: none"> ① Check the transducer position. ② Tighten the transducer and apply little coupling gel. ③ Ask the patient to walk slightly. ④ Ask the patient to walk in the prescriptive area.
The battery icon does not display when charging the battery.	<ul style="list-style-type: none"> ① The transducer does not connect to the charging point tightly. ② The base station is not supplied by AC power. 	<ul style="list-style-type: none"> ① Press the transducer to touch the charging point. ② Ensure the base station is supplied by AC power.
The charging board or charging point is corrosive.	<ul style="list-style-type: none"> ① It is wet or polluted by the coupling gel. 	<ul style="list-style-type: none"> ① Clean the transducer before charging. Replace the charging point if necessary.

B3.2 Blown Fuses

WARNING

Switch off the base station and remove the power cord before changing the fuse.

Replace the fuse when it is blown.

The two fuses of the base station are located on the rear panel, their specifications are:

Size: Φ5mm*20mm; Model: T2AH250V.

To replace a fuse:

- 1) Place the base station on a flat surface and remove the power cord.
- 2) Reverse the base station and pull the fuse container out as far as it can go.



- 3) Use a screw driver or a pair of pliers to push the fuse up from the bottom of the container.



- 4) Take the fuse out and replace it with a new one that is supplied by the manufacturer or of the same specifications.



- 5) Push the fuse container all the way back in position.

Appendix 4 Ultrasound Intensity and Safety

A4.1 Ultrasound in Medicine

The use of diagnostic ultrasound has proved to be a valuable tool in medical practice. Given its known benefits for non-invasive investigations and medical diagnosis, including investigation of the human fetus, the question of clinical safety with regards to ultrasound intensity arises.

There is no easy answer to the question of safety surrounding the use of diagnostic ultrasound equipment. Application of the ALARA (As Low As Reasonably Achievable) principle serves as a rule-of-thumb that will help you to get reasonable results with the lowest possible ultrasonic output.

The American Institute of Ultrasound in Medicine (AIUM) states that given its track record of over 25 years of use and no confirmed biological effects on patients or instrument operators, the benefits of the prudent use of diagnostic ultrasound clearly outweigh any risks.

A4.2 Ultrasound Safety and the ALARA Principle

Ultrasound waves dissipate energy in the form of heat and can therefore cause tissue warming. Although this effect is extremely low with Doppler, it is important to know how to control and limit patient exposure. Major governing bodies in ultrasound have issued statements to the effect that there are no known adverse effects from the use of diagnostic ultrasound, however, exposure levels should always be limited to As Low As Reasonably Achievable (the ALARA principle).

A4.3 Explanation of MI/TI

A4.3.1 MI (Mechanical Index)

Cavitations will be generated when ultrasound wave passes through and contacts tissues, resulting in instantaneous local overheating. This phenomenon is determined by acoustic pressure, spectrum, focus, transmission mode, and factors such as states and properties of the tissue and boundary. This mechanical bioeffect is a threshold phenomenon that occurs when a certain level of ultrasound output is exceeded. The threshold is related to the type of tissue. Although no confirmed adverse mechanical effects on patients or mammals caused by exposure at intensities typical of present diagnostic ultrasound instruments have ever been reported, the threshold for cavitation is still undetermined. Generally speaking, the higher the acoustic pressure, the greater the potential for mechanical bioeffects; the lower the acoustic frequency, the greater the potential for mechanical bioeffects.

The AIUM and NEMA formulate mechanical index (MI) in order to indicate the potential for mechanical effects. The MI is defined as the ratio of the peak-rarefactional acoustic pressure (should be calculated by tissue acoustic attenuation coefficient 0.3dB/cm/MHz) to the acoustic frequency.

$$MI = \frac{P_{r,\alpha}}{f_{awf} \times C_{MI}}$$

$C_{MI} = 1 \text{ (MPa / MHz)}$

A4.3.2 TI (Thermal Index)

Heating of tissues is caused by absorption of ultrasound when the ultrasound energy is applied. The temperature rise is determined by the acoustic intensity, exposed area and thermophysical properties of the tissue.

In order to indicate the potential for temperature rise caused by thermal effects, the AIUM and NEMA formulate thermal index (TI). It is defined as the ratio of the total acoustic power to the acoustic power required to raise the tissue temperature by 1 °C (1.8 °F).

According to different thermophysical properties of the tissue, TI is divided into three kinds: TIS, TIB and TIC.

TIS (Soft Tissue Thermal Index): It provides an estimate of potential temperature rise in soft or similar tissues.

TIB (Bone Thermal Index): It provides an estimate of potential temperature rise when the ultrasound beam passes through soft tissue and a focal region is in the immediate vicinity of bone.

TIC (Cranial Bone Thermal Index): It provides an estimate of potential temperature rise in the cranial bones or superficial bones.

A4.3.3 Measurement Uncertainty

The uncertainties in the measurements were predominantly systematic in origin; the random uncertainties were negligible in comparison. The overall systematic uncertainties were determined as follows.

1. Hydrophone Sensitivity

Based on the HNP-0400 hydrophone calibration certificate, the hydrophone measurement uncertainty for 1-15MHz is 1 dB, which is equivalent to an uncertainty of ±12.20% for intensity and ±6.10% for pressure. This uncertainty is used in PW measurement uncertainty assessment.

2. Digitizer

Based on the oscilloscope calibration certificate, the oscilloscope uncertainty is ±1.16% for intensity and ±0.58% for pressure.

3. Temperature

Based on the temperature variation of the water bath, the uncertainty is ±1.6% for intensity and ±0.8% for pressure.

4. Spatial Averaging

±10.2% for intensity, and ±6.1% for pressure.

5. Non-linear Distortion:

N/A. No effects of nonlinear propagation were observed.

Since all the above error sources are independent, they may be added on an RMS basis, giving a total uncertainty of ±26.62 percent for all intensity values reported, ±13.31 percent for all the pressure values and ±14.52 percent for the Mechanical Index.

A4.4 Prudent Use Statement

Although no confirmed bioeffects on patients caused by exposure from present diagnostic ultrasound equipment have ever been reported, the potential exists that such bioeffects may be identified in the future. Therefore, the ultrasound should be used prudently. High levels of acoustic output and long exposure time should be avoided while acquiring necessary clinical information.

A4.5 References for Acoustic Output and Safety

1. "Bioeffects and Safety of Diagnostic Ultrasound" issued by AIUM in 1993
2. "Medical Ultrasound Safety" issued by AIUM in 1994
3. "Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment, Revision 3" issued by AIUM/NEMA in 2004
4. "Standard for real-time display of thermal and mechanical acoustic output indices on diagnostic ultrasound equipment, Revision 2" issued by AIUM/NEMA in 2004
5. "Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers" issued in 2008.
6. "Medical electrical equipment—Part 2-37: Particular requirements for the basic safety and essential performance of ultrasonic medical diagnostic and monitoring equipment" issued by IEC in 2007.

A4.6 Transducer Acoustic Output Parameters List

A4.6.1 Test of Wired Transducer

Acoustic output reporting table for IEC60601-2-37

(IEC60601-2-37, Edition 2.1, 2015-06, table 201.103)

Operating Mode: PW mode

Working Frequency: 1.0MHz

Index label		MI	TIS		TIB		TIC
			At surface	Below surface	At surface	Below surface	
Maximum index value		0.013	0.0091		0.051		N/A
Index component value			N/A	0.0091	N/A	0.051	
Acoustic Parameters	$p_{r,a}$ at z_{MI} (MPa)	0.013					
	P (mW)		8.22		8.22		N/A
	P_{1x1} (mW)		N/A		N/A		
	z_s (cm)			13.10			
	z_b (cm)					13.10	
	z_{MI} (cm)	12.80					
	$z_{PPI,a}$ (cm)	12.80					
Other Information	f_{awf} (MHz)	1.00	1.00		1.00		N/A
	prr (Hz)	2000.00					
	srr (Hz)	N/A					
	n_{pps}	N/A					
	$I_{pa,a}$ at $z_{PPI,a}$ (W/cm^2)	0.011					
	$I_{spta,a}$ at $z_{PPI,a}$ or $z_{SII,a}$ (mW/cm^2)	1.92					
	I_{spta} at z_{PPI} or z_{SII} (mW/cm^2)	4.64					
Operating control conditions	p_r at z_{PPI} (MPa)	0.020					
	Focus(mm)	Fixed					
	Depth(mm)	Fixed					
	Frequency(MHz)	1.00					

Operating Mode: PW mode

Working Frequency: 1.0MHz

Acoustic Output		MI	$I_{SPTA,3}$ (mW/cm^2)	$I_{SPPA,3}$ (W/cm^2)
Global Maximum Value		0.013	1.92	0.011
Associated Acoustic Parameter	$P_{r,3}$ (MPa)	0.013		
	W_0 (mW)		8.22	8.22
	f_c (MHz)	1.00	1.00	1.00
	Z_{sp} (cm)	12.80	12.80	12.80
	Beam x-6 (cm)		0.62	0.62

	dimensions	y-6 (cm)		0.70	0.70
	PD (usec)		89.72		89.72
	PRF (Hz)		2000.00		2000.00
	EBD	A _z (cm)		Φ3.46	
Operating Control Conditions		E _{le} (cm)		Φ3.46	
		Focus(mm)		Fixed	
		Depth(mm)		Fixed	
		Frequency(MHz)		1.00	

A4.6.2 Test of Wireless Transducer (FTS-3)

A4.6.2.1 12-Crystal Wafer Transducer

Acoustic output reporting table for IEC60601-2-37

(IEC60601-2-37, Edition 2.1, 2015-06, table 201.103)

Operating Mode: PW mode

Working Frequency: 1.0MHz

Index label		MI	TIS		TIB		TIC
			At surface	Below surface	At surface	Below surface	
Maximum index value		0.017	0.0079		0.064		N/A
Index component value			N/A	0.0079	N/A	0.064	
Acoustic Parameters	p _{r,a} at z _{MI} (MPa)	0.017					
	P (mW)		9.69		9.69		N/A
	P _{1x1} (mW)		N/A		N/A		
	z _s (cm)			6.55			
	z _b (cm)					6.55	
	z _{MI} (cm)	6.55					
	z _{PII,a} (cm)	6.55					
	f _{awf} (MHz)	1.00	1.00		1.00		N/A
Other Information	prr (Hz)	2000.00					
	srr (Hz)	N/A					
	n _{pps}	N/A					
	I _{pa,a} at z _{PII,a} (W/cm ²)	0.0092					
	I _{spta,a} at z _{PII,a} or z _{SII,a} (mW/cm ²)	1.66					
	I _{spta} at z _{PII} or z _{SII} (mW/cm ²)	4.34					
	p _r at z _{PII} (MPa)	0.023					
Operating control conditions	Focus(mm)	Fixed					
	Depth(mm)	Fixed					
	Frequency(MHz)	1.00					

Operating Mode: PW mode
 Working Frequency: 1.0MHz

Acoustic Output		MI	$I_{SPTA,3}$ (mW/cm ²)	$I_{SPPA,3}$ (W/cm ²)
Global Maximum Value		0.017	1.66	0.0092
Associated Acoustic Parameter	$p_{r,3}$ (MPa)	0.017		
	W_0 (mW)		9.69	9.69
	f_c (MHz)	1.00	1.00	1.00
	z_{sp} (cm)	6.55	6.55	6.55
	Beam dimensions	x_{-6} (cm) y_{-6} (cm)	0.48 0.56	0.48 0.56
	PD	(usec)	90.07	
	PRF	(Hz)	2000.00	
	EBD	Az. (cm)	$\Phi 3.46$	
		Ele. (cm)	$\Phi 3.46$	
Operating Control Conditions	Focus(mm)		Fixed	
	Depth(mm)		Fixed	
	Frequency(MHz)		1.00	

A4.6.2.2 7-Crystal Wafer Transducer

**Acoustic output reporting table for IEC60601-2-37
(IEC60601-2-37, Edition 2.1, 2015-06, table 201.103)**

Operating Mode: PW mode
 Working Frequency: 1.0MHz

Index label		MI	TIS		TIB		TIC
			At surface	Below surface	At surface	Below surface	
Maximum index value		0.030	0.050		0.020		N/A
Index component value			N/A	0.050	N/A	0.020	
Acoustic Parameters	$p_{r,a}$ at z_{MI} (MPa)	0.030					
	P (mW)		9.83		9.83		N/A
	P_{1x1} (mW)		N/A		N/A		
	z_s (cm)			1.49			
	z_b (cm)					1.49	
	z_{MI} (cm)	1.49					
	$z_{PII,a}$ (cm)	1.49					
	f_{awf} (MHz)	1.00	1.00		1.00		N/A
Other Information	prr (Hz)	2000.00					
	srr (Hz)	N/A					
	n_{pps}	N/A					
	$I_{pa,a}$ at $z_{PII,a}$ (W/cm ²)	0.020					
	$I_{spta,a}$ at $z_{PII,a}$ or $z_{SII,a}$ (mW/cm ²)	3.17					
	I_{spta} at z_{PII} or z_{SII} (mW/cm ²)	3.59					

	p_r at z_{PII} (MPa)	0.030					
Operating control conditions	Focus(mm)	Fixed					
	Depth(mm)	Fixed					
	Frequency(MHz)	1.00					

Operating Mode: PW modeWorking Frequency: 1.0MHz

	Acoustic Output	MI	$I_{SPTA,3}$ (mW/cm ²)	$I_{SPPA,3}$ (W/cm ²)
	Global Maximum Value	0.030	3.17	0.020
Associated Acoustic Parameter	$p_{r,3}$ (MPa)	0.030		
	W_0 (mW)		9.83	9.83
	f_c (MHz)	1.00	1.00	1.00
	z_{sp} (cm)	1.49	1.49	1.49
	Beam dimensions	x ₋₆ (cm)	3.12	3.12
		y ₋₆ (cm)	3.12	3.12
	PD (usec)	91.26		91.26
	PRF (Hz)	2000.00		2000.00
	EBD	Az. (cm)	Φ 3.12	
		Ele. (cm)	Φ 3.12	
Operating Control Conditions	Focus(mm)		Fixed	
	Depth(mm)		Fixed	
	Frequency(MHz)		1.00	

A4.6.3 Standard Parameter Equal Contrast List

IEC60601-2-37 standard parameter equal contrast list	
IEC60601-2-37 parameter	NOTE
$p_{r,a}$	Attenuated Peak-rare-factional Acoustic Pressure
p_r	Peak-rare-fractional Acoustic Pressure
P	Output Power
z_s	Depth for Soft Tissue Thermal Index
$P_a(Z_s)$	Attenuated Output Power
$I_{ta,a}(Z_s)$	Attenuated Temporal-average Intensity
z_{bp}	Break-point Depth
z_b	Depth for Bone Thermal Index
$I_{pi,a}$	Attenuated Pulse-intensity Integral
I_{pi}	Pulse-intensity Integral
$d_{eq}(Z_b)$	Equivalent Beam Diameter at the point of Z_{sp}
f_{awf}	Center Frequency, Acoustic Working Frequency
X	-12dB Output Beam Dimensions
Y	
t_d	Pulse Duration
prr	Pulse Repetition Frequency (Pulse Repetition Rate)
d_{eq}	Equivalent Beam Diameter
FL_x	Focal Length
FL_y	
$I_{pi,a}$ at max MI	Attenuated Pulse-average Intensity at the point of Maximum MI
A_{aprt}	-12dB Output Beam Area
MI	Mechanical Index
TIS	Soft Tissue Thermal Index
TIB	Bone Thermal Index
TIC	Cranial-bone Thermal Index

parameter specified in TRACK1 of FDA Guidance	
TRACK1 parameter	NOTE
p _{r.3}	Derated Peak-rare-factional Acoustic Pressure
W ₀	Output Power
Z _{sp}	Z _{sp} = Z _{B.3} , Depth for Bone Thermal Index
f _c	Center Frequency, Acoustic
X ₋₆	-6dB Beamwidth
y ₋₆	
PD	Pulse Duration
PRF	Pulse Repetition Frequency
MI	Mechanical Index
I _{SPTA.3}	Derated Spatial-peak Temporal-average Intensity
I _{SPPA.3}	Derated Spatial-peak Pulse-average Intensity
Az.	Aperture X width Y Dimeter
Ele.	
EDS	Entrance Dimensions Of The Scan
EBD	Entrance Beam Dimensions

Appendix 5 Abbreviation

The abbreviations used in this manual and their full names are listed below:

Abbreviation	Full Name
AC	Alternative Current
AFM	Automatic Fetal Movement [Detection]
BPM	Beat(s) Per Minute
CTG	Cardiotocography
DC	Direct Current
DECG	Direct ECG
DFHR	Direct FHR
DIA	Diastolic Blood Pressure
ECG	Electrocardiogram
FH	Fetal Heart
FHR	Fetal Heart Rate
FM	Fetal Movement
FS	Fetal Stimulator
MHR	Maternal Heart Rate
ICU	Intensive Care Unit
ID	Identity
IUP	Intra-Uterine Pressure
IUPC	Intra-Uterine Pressure Catheter
LCD	Liquid Crystal Display
MAP	Mean Artery Blood Pressure
MECG	Maternal ECG
MFM	Manual Fetal Movement [Detection]
MRI	Magnetic Resonance Imaging
NIBP	Non-Invasive Blood Pressure
NST	Non Stress Test
PR	Pulse Rate
RF	Radio Frequency
SOV	Signals Overlap Verification
SpO ₂	Pulse Oximetry

STV	Short-Term Variation
SYS	Systolic Blood Pressure
TEMP	Temperature
TOCO	Tocotonometer
UA	Uterine Activity [TOCO/IUP]
US	Ultrasound [Transducer]

Appendix 6 Ordering Information

Accessories (standard and optional configuration) supplied or approved by the manufacturer can be used with the monitors. See the following table for details.

Part Number	Accessory	Specification
02.01.107705	US Transducer	12 ultrasound crystals, 1MHz, yellow label
02.01.31528	US Transducer	12 ultrasound crystals, 1MHz, purple label
02.01.210821	US-T Transducer	12 ultrasound crystals, 1MHz, IEC Standard
02.01.212717	US-T Transducer	7 ultrasound crystals, 1MHz, IEC Standard
02.01.31527	TOCO Transducer	Blue label
02.01.107791	TOCO Transducer	Cyan label
02.01.210823	TOCO-T Transducer	IEC Standard
02.01.211642	TOCO-E Transducer	IEC Standard
01.13.104152	IUP Cable	PN:56321
01.57.02145	Disposable Fetal Spiral Electrode	PN:31479549
01.57.02146	Disposable Maternal Attachment Pad Electrode	PN:50000095
01.57.104153	Disposable Intrauterine Pressure Catheter	PN:56300
01.13.036358	DECG Cable	TPU,L=2200mm
01.13.036477	DECG Cable	TPU,L=2200mm DECG-Q
01.13.036478	DECG Cable	TPU,L=2200mm DECG-P
01.57.471610	FTS-3 DECG Cable	DECG-WT
01.57.471608	FTS-3 DECG Cable	DECG-WQ
01.57.471609	FTS-3 DECG Cable	DECG-WP
01.13.036357	IUP Connecting Cable	L=150mm
02.01.210095	Remote Event Marker	/2.5m
01.57.471447	Belt	/1400mm*58mm
02.06.17661	Fetal Stimulator	
01.57.78001	Ultrasound Gel	/PARKER
01.57.78008-11	Aquasonic Coupling Gel	/Shenfeng
01.57.471095	3-lead ECG Cable	AHA Standard, Grabber Style
01.57.471098	3-lead ECG Cable	IEC Standard, Grabber Style
01.57.471383	3-lead ECG Cable	IEC Standard, Grabber Style
01.57.471384	3-lead ECG Cable	AHA Standard, Grabber Style
01.57.471501	FTS-3 MECG Cable	AHA Standard, Snap Style
01.57.471503	FTS-3 MECG Cable	AHA Standard, Grabber Style
01.57.471502	FTS-3 MECG Cable	IEC Standard, Snap Style
01.57.471504	FTS-3 MECG Cable	IEC Standard, Grabber Style

01.57.471276	Disposable ECG Electrode	For Adult
02.01.109069	SpO2 Sensor	/SH1
01.15.030043	SpO2 Sensor	NELLCOR
01.57.471005	NIBP Cuff Extension Tube	/TPU, Grey
01.57.471330	NIBP Cuff	For Adult, 27-35cm,E9
01.57.471331	NIBP Cuff	For Large Adult, 34-43cm,E10
01.15.040420	TEMP Sensor	Skin Contact
01.15.30043	SpO2 Sensor	NELLCOR / DS-100A
01.13.30131	SpO2 Sensor Extension cable	NELLCOR / DS-100A
01.57.75111	Thermosensitive Paper	GE, AHA Standard, with green safe range
01.57.471047	Thermosensitive Paper	GE, AHA Standard, without green safe range
01.57.75112	Thermosensitive Paper	GE, IEC Standard
01.57.75113	Thermosensitive Paper	Phillips, AHA Standard
01.57.75114	Thermosensitive Paper	Phillips, IEC Standard
01.13.037122	Power Cord	AHA Standard
01.13.36014	Power Cord	IEC Standard
21.21.064150	Rechargeable Lithium-ion Battery	5000 mAh
21.21.064181	Fuse	T2AH250V
01.13.036770	Signal Cable	USB TO DB9
01.13.20096	Signal Cable	Downloading cable/ Ethernet
01.13.036124	Signal Cable	F9 to OB TraceVue cable (DB9 to RJ45, For OB TraceVue with adapter)
01.13.107974	Signal Cable	F9 to OB TraceVue cable (DB9 to RJ45)
01.13.107702	Signal Cable	F9 to OB TraceVue cable (DB9 to DB9)
02.01.210517	Wireless AP module	WL-330N,for F3/F6/F9
01.18.052246	Wireless AP	DWL-3200AP
01.57.471026	Dust Cover	890mm*740mm
01.55.461211	Keypad Cover	assembly for F9
01.24.070019	Screwdriver	Φ 4
01.13.114214	Potential Equalization Conductor	Foreign Sales
01.13.036299	Signal Cable	Wireless Transducer Connecting Cable
01.13.036301	Signal Cable	Wireless Transducer Y Shape Connecting Cable
01.21.064143	Rechargeable Lithium-ion Battery	5000 mAh
02.04.240301	Tablet	MT-803
02.01.210926	Basket	MT-803
83.62.108166	Trolley	MT-503N plus laptop holder, provided with F9
83.60.108108	Trolley	MT-503, provided with F9
02.01.108125	Adapter Plate	Provided with F9
02.04.101976	Basket	MT-503/MT-206
02.01.112910	Wall Mounting Package	Provided with F9
02.01.112891	Basket	Provided with F9/ Wall Mounting

02.01.210061	Wall-Mounting Bracket	Provided with F9/ With basket
01.56.465631	Carry Bag	
	Quick Reference	
	User Manual	
	Service Manual	
	Fetal Heart Monitoring Guidance	

The accessories employed by the manufacturer, such as the rechargeable battery, are products having passed the authentication of CE, and they have the characteristics specified by their manufacturers. The materials with which the patient can come into contact conform to the standard of ISO 10993.

CAUTION

Replacement of all above accessories can be performed by the operator. But only the accessories supplied or recommended by the manufacturer are allowed connected to the monitor and FTS-3 system.

Appendix 7 EMC Information

A7.1 Electromagnetic Emissions

Guidance and manufacturer's declaration – electromagnetic emission		
The <i>F9 and F9 Express Fetal & Maternal Monitors</i> are intended for use in the electromagnetic environment specified below. The customer of the user of the <i>F9 and F9 Express Fetal & Maternal Monitors</i> should assure that it is used in such and environment.		
Emission test	Compliance	Electromagnetic environment – guidance
RF emissions CISPR 11	Group 1	The <i>F9 and F9 Express Fetal & Maternal Monitors</i> use RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emission CISPR 11	Class A	The <i>F9 and F9 Express Fetal & Maternal Monitors</i> are suitable for use in all establishments, other than domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Harmonic emissions IEC 61000-3-2	Class A	
Voltage fluctuations/ flicker emissions IEC 61000-3-3	Complies	

A7.2 Electromagnetic Immunity

Guidance and manufacturer's declaration – electromagnetic immunity			
The <i>F9 and F9 Express Fetal & Maternal Monitors</i> are intended for use in the electromagnetic environment specified below. The customer or the user of <i>F9 and F9 Express Fetal & Maternal Monitors</i> should assure that it is used in such an environment.			
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Electrostatic discharge (ESD) IEC 61000-4-2	± 8 kV contact ±15 kV air	± 8 kV contact ± 15 kV air	Floors should be wood, concrete or ceramic tile. If floor are covered with synthetic material, the relative humidity should be at least 30%.
Electrical fast transient/burst IEC 61000-4-4	± 2 kV for power supply lines ± 1 kV for input/output lines	± 2kV for power supply lines Not applicable	Mains power quality should be that of a typical commercial or hospital environment.
Surge IEC 61000-4-5	± 1 kV line(s) to line(s) ± 2 kV line(s) to ground	± 1 kV line(s) to line(s) ± 2 kV line(s) to ground	Mains power quality should be that of a typical commercial or hospital environment.
Power frequency (50Hz/60Hz) magnetic field IEC61000-4-8	30 A/m	30 A/m	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.

<p>Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11</p>	<p>0 % U_T; 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°</p> <p>0 % U_T; 1 cycle and 70 % U_T; 25/30 cycles) Single phase: at 0°</p> <p>0 % U_T; 250/300 cycle</p>	<p>0 % U_T; 0,5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315°</p> <p>0 % U_T; 1 cycle and 70 % U_T; 25/30 cycles) Single phase: at 0°</p> <p>0 % U_T; 250/300 cycle</p>	<p>Mains power quality should be that of a typical commercial or hospital environment. If the user of the <i>F9 and F9 Express Fetal & Maternal Monitors</i> requires continued operation during power mains interruptions, it is recommended that the <i>F9 and F9 Express Fetal & Maternal Monitors</i> be powered from an uninterruptible power supply or a battery.</p>
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NOTE: U_T is the a.c. mains voltage prior to application of the test level.

A7.3 Electromagnetic Immunity

Guidance and manufacture's declaration – electromagnetic immunity			
The <i>F9 and F9 Express Fetal & Maternal Monitors</i> are intended for use in the electromagnetic environment specified below. The customer or the user of <i>F9 and F9 Express Fetal & Maternal Monitors</i> should assure that it is used in such an environment.			
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance
Conducted RF IEC 61000-4-6	3 Vrms 150 kHz to 80 MHz 6VrmSC)in ISM bands between 0,15 MHz and 80 MHz	3Vrms 6VrmSC)in ISM bands between 0,15 MHz and 80 MHz	<p>Portable and mobile RF communications equipment should be used no closer to any part of the <i>F9 and F9 Express Fetal & Maternal Monitors</i> including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.</p> <p>Recommended separation distance</p> $d = 1.2\sqrt{P} \quad 150 \text{ kHz to } 80 \text{ MHz}$ $d = 1.2\sqrt{P} \quad 80 \text{ MHz to } 800 \text{ MHz}$ $d = 2.3\sqrt{P} \quad 800 \text{ MHz to } 2.7 \text{ GHz}$ <p>$d = 6\sqrt{P} / E$ at RF wireless communications equipment bands (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 monitor, including cables specified by the manufacturer).</p> <p>Where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer and d is the recommended separation distance in meters (m).</p> <p>Field strengths from fixed RF transmitters, as determined by an electromagnetic site</p>
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.7 GHz	3 V/m 80 MHz to 2.7 GHz	

			<p>survey,^{a)} should be less than the compliance level in each frequency range.^{b)}</p> <p>Interference may occur in the vicinity of equipment marked with the following symbol:</p> 
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.			
<p>a) Field strengths from fixed transmitters, such as base stations for radio (cellular/wireless) 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 the <i>F9 and F9 Express Fetal & Maternal Monitors</i> are used exceeds the applicable RF compliance level above, the <i>F9 and F9 Express Fetal & Maternal Monitors</i> should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the <i>F9 and F9 Express Fetal & Maternal Monitors</i>.</p> <p>b) Over the frequency range 150 kHz to 80 MHz, field strengths should be less than 3 V/m.</p> <p>c) The ISM (industrial, scientific and medical) bands between 0.15 MHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz. The amateur radio bands between 0.15 MHz and 80 MHz are 1.8 MHz to 2.0 MHz, 3.5 MHz to 4.0 MHz, 5.3 MHz to 5.4 MHz, 7 MHz to 7.3 MHz, 10.1 MHz to 10.15 MHz, 14 MHz to 14.2 MHz, 18.07 MHz to 18.17 MHz, 21.0 MHz to 21.4 MHz, 24.89 MHz to 24.99 MHz, 28.0 MHz to 29.7 MHz and 50.0 MHz to 54.0 MHz.</p>			

Table-Test specifications for ENCLOSURE PORT IMMUNITY to RF wireless communications equipment

Test Frequency (MHz)	Brand ^{a)} (MHz)	Service ^{a)}	Modulation ^{b)}	Maximum Power(W)	Distance (m)	IMMUNITY TEST LEVEL (V/m)
385	380-390	TETRA 400	Pulse modulation ^{b)} 18Hz	1.8	0.3	27
450	430-470	GMRS 460, FRS 460	FM ^{c)} ±5 kHz deviation 1kHz sine	2	0.3	28
710	704-787	LTE Brand 13, 17	Pulse modulation ^{b)} 217 Hz	0.2	0.3	9
745						
780						
810						
870	800-960	GSM 800/900,TETRA 800, iDEN 820, CDMA 850, LTE	Pulse modulation ^{b)} 18 Hz	2	0.3	28
930						

		Band 5				
1720	1700-1990	GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4,25; UMTS	Pulse modulation ^{b)} 217 Hz	2	0.3	28
1845						
1970						
2450	2400-2570	Bluetooth, WLAN,802.11 b/g/n, RFID 2450, LTE Brand 7	Pulse modulation ^{b)} 217 Hz	2	0.3	28
5240	5100-5800	WLAN 802.11 a/n	Pulse modulation ^{b)} 217 Hz	0.2	0.3	9
5500						
5785						
<p>Note: If necessary to achieve the IMMUNITY TEST LEVEL, the distance between the transmitting antenna and the ME EQUIPMENT or ME SYSTEM maybe reduce to 1m. The 1 m test distance is permitted by IEC 61000-4-3.</p> <p>a) For some services, only the uplink frequencies are included.</p> <p>b) The carrier shall be modulated using a 50% duty cycle square wave signal.</p> <p>c) As an alternative FM modulation, 50% pulse modulation at 18 Hz may be used because while it does not represent actual modulation, it would be worst case</p>						

A7.4 Recommended Separation Distances

**Recommended separation distances between
portable and mobile RF communications equipment and the
F9 and F9 Express Fetal & Maternal Monitors**

The *F9 and F9 Express Fetal & Maternal Monitors* are intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the *F9 and F9 Express Fetal & Maternal Monitors* can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the *F9 and F9 Express Fetal & Maternal Monitors* as recommended below, according to the maximum output power of the communications equipment.

Rated maximum output power of transmitter (W)	Separation distance according to frequency of transmitter (m)		
	150 kHz to 80 MHz $d = 1.2\sqrt{P}$	80 MHz to 800 MHz $d = 1.2\sqrt{P}$	800 MHz to 2.7 GHz $d = 2.3\sqrt{P}$
0.01	0.12	0.12	0.23
0.1	0.38	0.38	0.73
1	1.2	1.2	2.3
10	3.8	3.8	7.3
100	12	12	23

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

NOTE 1: At 80 MHz and 800 MHz, the separation distance for 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.

Appendix 8 Limitations of Ultrasonic Monitoring

A8.1 How Does Ultrasound Work

When the ultrasound waves strike an object, they bounce back and create an echo. If the object moves toward the sound source, the frequency of the echo increases. If the object moves away from the sound source, the frequency of the echo decreases. This is called “Doppler Effect”. In the 1960's, the ultrasonic technique was first applied to medical diagnostic imaging.

The ultrasound process involves placing a small device called a transducer, against the skin of the patient near the region of interest. The ultrasound transducer combines functions of emitting and receiving ultrasounds in one device. This transducer produces a stream of inaudible, high frequency sound waves which penetrate into the body and bounce off the organs inside. It detects sound waves as they bounce off or echo back from the internal structures and contours of the organs. The movement of the organs produces the Doppler Effect, and this movement can be measured and described by measuring the echo.

In fetal monitoring, the ultrasound transducer produces a stream of sound waves which penetrate into the maternal abdomen and bounce off the fetal heart. Then the transducer receives the echoes and transfers them to the monitor, which turns the signal into fetal heart beating sound and fetal heart rate trace.

Therefore, placement of the transducer is critical to ultrasound fetal heart monitoring.

A8.2 Artifacts in Fetal Heart Monitoring

(1) How does artifact happen?

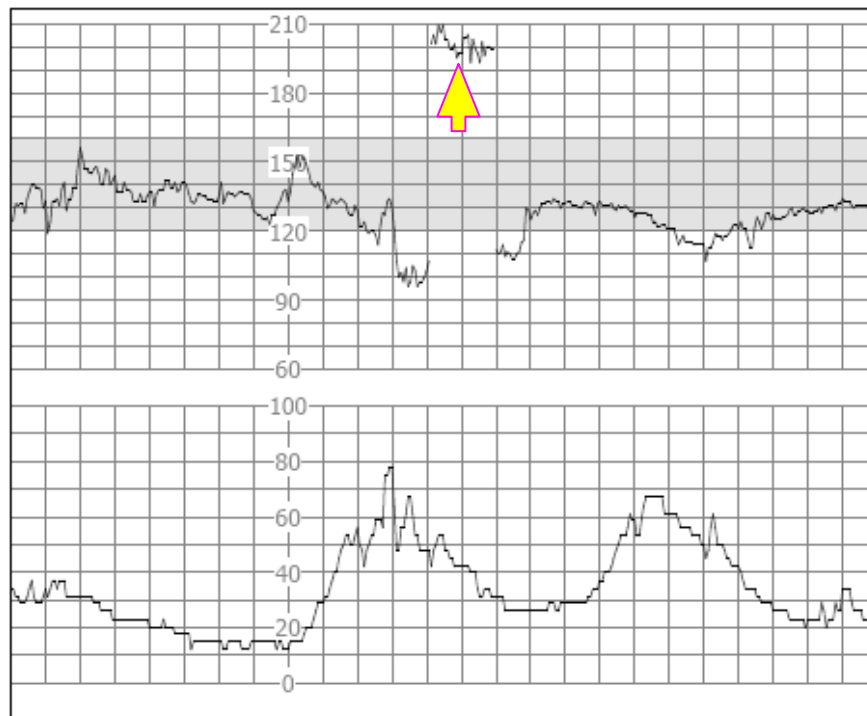
The transducer detects sound waves as they bounce off or echo back from the fetal heart. However, the sound waves bouncing off from maternal blood vessels may be detected by the transducer and then be processed by the monitor as well. As a result, artifacts may be produced.

The artifacts, if not correctly interpreted, may cause the physicians to perform unnecessary interventions, or to fail to detect the fetal distress and the need for interventions.

The most common artifacts are doubling and halving.

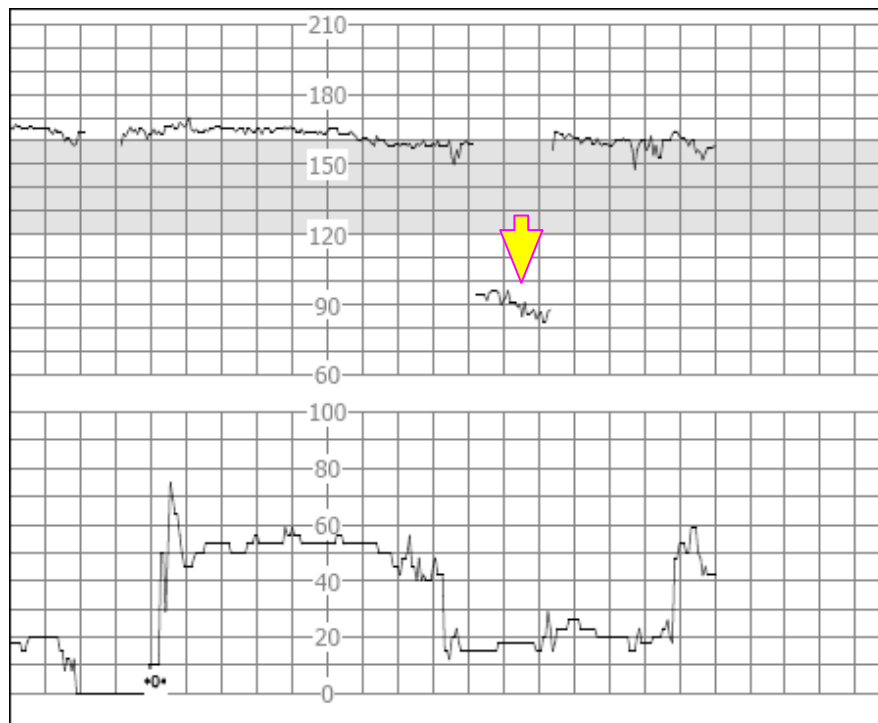
(2) Doubling:

When the FHR drops to 120 bpm or lower, the diastole and systole become far apart, thereby the monitor may mistake these two movements of a single heartbeat for two separate heartbeats. As a result, a heart rate trace that is double the actual heart rate is produced. This often happens during severe decelerations and bradycardia, representing an abrupt switch of the trace to double the actual heart rate.



(3) Halving:

When the FHR increases to 180 bpm or higher, it is possible for the monitor to mistake the two separate hearbeats for the diastole and systole of a single heartbeat. As a result, a heart rate trace that is half the actual heart rate is produced. This often happens during tachycardia, representing an abrupt switch of the trace to half the actual heart rate. The clinicians may interpret it as a “deceleration”.



However, the heart beat sound from the monitor speaker is still reliable even when doubling or halving is occurring.

Stethoscopy should be applied when sudden changes in baseline are detected.

If the amniotic membrane rupture and cervical dilatation are sufficient, consider using a spiral electrode to obtain precise FHR with direct fetal ECG as the signal source.

(4) Erratic Traces / Drop out

When the fetal heart moves partially out of the ultrasound wave path, the transducer receives mixed or weak signals, and thereby the monitor presents erratic traces. When the fetal heart moves fully out of the path, inadequate consecutive and periodic signals are received, and no trace is represented.

Erratic traces and transitory episodes of drop out are common, especially when the fetus or/and mother move(s). If they exist for an extended period, it indicates that the transducer is not aimed at the fetus. Repositioning of the transducer is needed.

A8.3 Audio Output and Screen Reading

In most instances, the audio output from the monitor speaker corresponds to the readings presented on the monitor screen. But occasionally the fetal heart sound may differ from the trace and numeric.

When the fetal heart moves partially out of the ultrasound wave path, the transducer receives weaker FHR signal and other stronger signals (usually maternal heart/pulse rate). After the signals are transmitted to the monitor, the audio system and the video system of the monitor process the signals separately. On one hand, the audio circuit filters the low-frequency signals and gives audio output of the high-frequency signals, so fetal heart sound is heard. On the other hand, the autocorrelation algorithm computes the stronger signal source and thereby the maternal heart/pulse rate is displayed. As a result, the audio output differs from the screen reading.

If this situation occurs, it can be dismissed by repositioning the transducer.

In a word, the abnormalities listed above (artifacts, sound and reading differences) are caused by the limitations of ultrasonic monitoring technique. Fortunately they rarely occur. But a good understanding of how to detect them and what countermeasures should be taken will help obtain better fetal monitoring effect.

We hope you find this information useful. If you have any questions about fetal monitoring, please contact our sales representatives and perinatal specialists.

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