

JUMPER

User Manual

Fetal Doppler

Model: SHA20

Manual Version: 3.0
Issuing Date: 2024.4

Thank you for purchasing the Fetal Doppler made by Jumper Medical. Before using the product, read this manual carefully and operate the product as specified in this manual.

SECTION 1: INTRODUCTION

1.1 PACKING LIST

Main unit × 1;
Battery AA × 2;
Coupling agent × 1 (Optional);
User manual × 1.

1.2 PRODUCT DESCRIPTION

The product is mainly used to detect the sound of the fetal heartbeat (SFH). Fetal heart rate (FHR) is an important basis for checking whether a fetus is healthy. Recording the changes in FHR helps detect signs of fetal hypoxia, fetal distress, fetal umbilical cord around the neck, and so on. Fetal monitoring at home mainly includes listening to fetal heartbeat and checking FHR changes, which help greatly increase fertility safety.

1.3 OPERATING PRINCIPLE

Based on the Doppler principle, a 3.0MHz ultrasonic probe is used to capture fetal heart signals from the belly of a pregnant woman. After signal processing of the backend circuit, fetal heart signals are output to speakers to play sound. Audio signals are wirelessly sent by using the built-in Bluetooth module. A smartphone that has connected to the product receives the data and calculates and displays fetal heart rate information by using specified mobile phone software.

FCC ID: 2ADYL-SHA20

SECTION 2: SAFETY GUIDANCE

2.1 INTENDED USE

The Fetal Doppler SHA20 is a hand-held, battery-powered audio Doppler device used for detecting fetal heartbeats. The patient is an intended operator.

2.2 INDICATIONS FOR USE

The product is normally applied to the fetus above 12 weeks of gestation, difference in pregnant women.

- Listen to SFH: Operator can listen to the sound of fetal heartbeat from the speaker.
- Audio record: The sound of fetal heartbeat can be recorded by APP.

CAUTION: It should not be used in life supporting or life-sustaining applications.

2.3 CONTRAINDICATIONS FOR USE

The device has no side-effects if administered correctly and residual risk is acceptable.

2.4 NOTE FOR HOME USE

This device cannot replace a professional fetal monitor. If the fetal heart rate is abnormal or cannot be located by using this monitor, pregnant women should immediately go to the hospital to seek the doctor's help. If fetal movement is not felt by the pregnant woman, immediately go to the hospital to seek the doctor's help.

2.5 SAFETY TERMS AND CONDITIONS

The signal words shown below left identify the potential hazard categories. The definition of each category is as follows:

DANGER: This alert identifies hazards that will cause serious personal injury or death.

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

CAUTION: This alert identifies hazards that may cause minor personal injury, product damage, or property damage.

2.6 SAFETY ALERT DESCRIPTIONS

The following is a list of product safety alerts that appear in this section and throughout this manual. You must read, understand, and pay heed to these safety alerts before attempting to operate the product.

DANGER: Fire and Explosion Hazard

Do not operate the product in the presence of flammable gases to avoid possible explosion or fire hazard.

WARNING: Strangulation resulting from baby or child entanglement in monitoring cables.

WARNING: Do not modify this equipment without authorization of the manufacturer.

WARNING: Dust, light may affect the safety and performance of the instrument.

WARNING: Degraded sensors and electrodes, or loosened electrodes, that can degrade performance or cause other problems.

WARNING: The effects caused by pets, pests or children

WARNING: Use only Approved Equipment

Do not use batteries, gel, cables, or optional equipment other than those approved by manufacturer which may cause the product to function improperly during a rescue.

WARNING: Adjacent and/or Stacked Equipment

The product should not be used adjacent to or stacked with other equipment. If adjacent or stacked use is necessary, the product should be observed to verify normal operation in the configuration in which it will be used.

CAUTION: Check that the equipment does not have visible evidence of damage that may affect personnel's safety or examining capability before use. If damage is detected, replacement is recommended.

CAUTION: The surface of the probe in contact with the patient may cause discomfort due to biocompatibility issues. The coupling agent may cause skin allergies in users. If the patient experiences any discomfort or allergic reactions, usage should be immediately discontinued and medical attention sought if necessary.

CAUTION: Do not wrap the probe wire to avoid suffocation.

CAUTION: Don't touch patient, power port, and probe at the same time.

CAUTION: This product is not recommended for use on ships and aircraft.

CAUTION: Please keep the Fetal Doppler and batteries out of the reach of children to prevent them from playing with them. In the event that a child accidentally swallows a battery, seek immediate medical attention.

CAUTION: Temperature/Humidity/Pressure extremes

Exposing the product to extreme environmental conditions outside of its operating parameters may compromise the ability of the product to function properly.

CAUTION: Battery Disposal

Recycle or dispose of the battery in accordance with the local laws. To avoid fire and explosion hazard, do not burn or incinerate the battery.

CAUTION: Possible Radio Frequency (RF) Susceptibility

RF susceptibility from cellular phones, CB radios and FM 2-way radio may cause interference with the product. Do not operate wireless radiotelephones in the vicinity of the product – turn power off to the radiotelephone and other like equipment near the product.

CAUTION: Systems Statement

Equipment connected to the product must be certified to the respective IEC Standards (IEC 60601-1 for medical equipment).

CAUTION: Case Cleaning Solutions

When disinfecting the case, use a non-oxidizing disinfectant, such as ammonium salts or glutaraldehyde-based cleaning solution, to avoid damage to the metal connectors.

CAUTION: Environment of use

The product is designed for indoor use. Operator must confirm that the environment of use meets the required operating environmental specifications before using.

CAUTION: Cold Environments

If the product is stored in an environment with a temperature below the operating temperature, the unit should be allowed to warm up to the needed operating temperature before using.

CAUTION: Do not use the device with HF surgical equipment.

2.7 SYMBOL DESCRIPTIONS

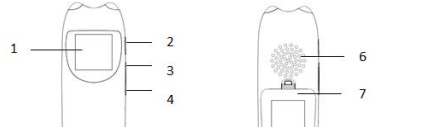
The following symbols may appear in this manual, on the product, or on its accessories. Some of the symbols represent standards and compliances associated with the product and its use.

| | |
|--|---|
| | Consult instructions for use of the product and/or its accessories. |
| | Medical Device |
| | Warning Information. |
| | Authorized Representative in the European Community. |
| | This product complies with the Regulation (EU) 2017/745 requirements. |
| | Date of manufacture. |
| | Manufacturer information. |
| | Type BF applied part |
| | Specifies serial number of the Product |
| | Batch code |
| | The environmental protection use period is 5 years. |
| | Degree of protection against ingress of water and particulate matter. |
| | It indicates that the equipment should be sent to the special agencies according to local regulation for separate collection after its useful life. |
| | Storage Temperature |
| | Humidity |
| | Atmospheric Pressure |

SECTION 3: USING THE PRODUCT

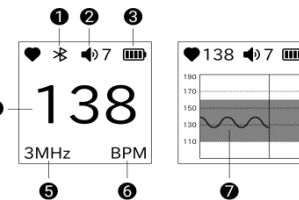
This section provides the description for operation.

3.1 PRODUCT STRUCTURE



1: TFT Display 2: Power/Mode button
3: Volume up button 4: Volume down button
5: Ultrasonic probe 6: Speaker
7: Battery compartments

3.2 INTERFACE DISPLAY



1: Bluetooth 2: Volume
3: Battery indicator 4: FHR
5: Ultrasonic frequency 6: Beat per minute
7: The curve of FHR

3.2.1 Power on/off

Power on: Press and hold button for about 2s and the screen lights up, and the device is powered on. In power on state, short press button to switch the display mode.

Power off: In the power-on state, press and hold button for about 2s, the screen goes out, and the device is powered off.

3.2.2 Volume adjustment button

While monitoring, the volume increases by press "+" button, and the volume decreases by press "-" button. There are 7 volume levels.

3.2.3 Icons

: In curve mode, when a heartbeat is detected, the fetal heartbeat symbol will light up and flash with the heart rate. In value mode, this symbol signifies the quality of the detected fetal heart signal. Three stars represent the best signal quality, while zero stars indicate that no fetal heartbeat signal was detected. It is important to follow the correct method to locate the optimal fetal heart position.

: The number next to the volume symbol indicates the volume and can be adjusted from 0-7.

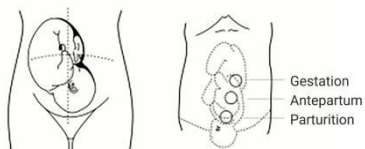
: When the icon displays in red , it means that the battery is about to run out and needs to be replaced in time.

: The Bluetooth icon is green when the Bluetooth is connected, and is white when disconnected.

3.3 USING PRODUCT TO DETECT

Locate the position of the fetus by hand touching, firstly to find out the best direction to the fetal heart. Place the faceplate of probe at the best position for detecting fetal heartbeat. Adjust the transducer to obtain an optimum audio signal ideally by angling the transducer around. Generally, the site of heart of fetus is 1/3 below the navel line at its earlier stage, it then moves upward with increasing of gestational period, and the site of heart of fetus will be a little deviation to left or right with different fetuses. Please make sure that the surface of the probe should be fully contacted with the skin. After the sound becomes clear, it is the proper functioning. If no coupling gel, water can be used.

Note: The normal range of fetal heart rate is 110 bpm-160 bpm. During measurement, values are displayed in white within the normal range and in red when they are out of range.



3.4 CONNECTING THE INSTRUMENT TO THE SMARTPHONE VIA BLUETOOTH

Software Downloading:

1. Download and install the mobile phone APP software "JUMPER Health" by scanning the QR code on the packing box or searching for the APP in application stores such as APP Store/Google Play.

2. This software supports iOS 7.0 and later versions, and Android 4.3 and later versions. In addition, hardware of the smartphone needs to support Bluetooth 4.0.

Bluetooth connection:

Start "JUMPER Health" on the smartphone, turn on the smartphone's Bluetooth function to search for the Bluetooth signal and pair the device.

Software Usage:

Detailed see software operation manual of JUMPER Health.

SECTION 4: MAINTENANCE & CLEANING AND DISINFECTION

4.1 MAINTENANCE

4.1.1 The transducer acoustic surface is fragile and must be handled with care. Gel must be wiped off from the transducer after use. These precautions will prolong the life of the unit.

4.1.2 To ensure the product is always functional when required, the following maintenance shall be performed.

- Visual Inspection
- Clean the product and its accessories
- Check the battery fuel gauge
- Test product performance
- Remove the battery if it is not used for a long time.
- The product requires no calibration.

Note: No service and maintenance while the equipment is in use. Manufacturer will make available on request circuit diagrams, component part lists, descriptions, calibration instructions, or other information that will assist to service personnel in parts repair.

4.2 CLEANING PRODUCT AND ACCESSORIES

The following cleaning products may be used to clean the exterior surfaces of the product.

- Isopropyl alcohol (70% solution in water)
- Mild soap and water
- Sodium hypochlorite (chlorine bleach) (3% solution in water).
- Quaternary ammonium compounds (such as Lysol) (10% solution in water).

WARNING: Do not use abrasive cleaners or strong solvents such as acetone or acetone-based cleaners.

WARNING: Do not use mixing disinfecting solutions (such as bleach and ammonia) as hazardous gases may result.

WARNING: Do not use acid, alkaline, or corrosive detergent.

WARNING: Do not clean electrical contacts or connectors with bleach.

4.3 CLEANING INSTRUCTIONS

1. Before cleaning the product, turn off the product.
2. Before cleaning, remove all adherent soil (tissue, fluids, etc.) and wipe thoroughly with a cloth dampened with water before applying the cleaning solution.
3. When cleaning, do not immerse. Keep the exterior surface of the device clean and free of dust and dirt, clean exterior surface of the unit with a dry, soft cloth, if necessary, clean it with a soft cloth soaked in a solution of soap and wipe dry with a clean cloth immediately. Wipe the transducer body with soft cloth to remove any remaining coupling gel. Clean with soap only.

CAUTION: To prevent damage to the product, do not clean any part of the Product or Accessories with phenolic compounds. Do not use abrasive or flammable cleaning agents. Do not steam, autoclave, or gas-sterilize the Product or accessories.

CAUTION: Cleaning liquids: do not submerge the product in liquids or pour cleaning liquids over, into or onto the product.

4.4 DISINFECTION

Cleaning the unit surface and the transducer as the above mentioned, then wipe the surface of transducer with 75% ethanol or alcohol, clean the transducer surface with a dry, soft cloth.

WARNING: Don't use low temperature steam sterilization or other way to sterilize.

WARNING: Don't use high temperature sterilizing process.

After cleaning or disinfection, check if the Doppler function well. If any problem is detected, please contact the manufacturer for service before reusing them.

Visual Check: Check if the Doppler probe and host are damaged:

Function Check:

1. Check if the Doppler can be switched on or off properly;

2. Check if the TFT works normally;

3. Rub the surface of the probe with your hand to check if the Doppler is producing sound properly

SECTION 5: SPECIFICATIONS & TROUBLESHOOTING

This section presents the specifications and safety standards of the Product.

5.1 SPECIFICATIONS

 Note: The following specifications are subject to change and are only noted as a point of reference.

Technical Specifications

| |
|--|
| Acoustic working frequency: 3.0MHz±5% |
| Overall sensitivity (200 mm off the probe surface): not lower than90 dB |
| Spatial-peak temporal-peak acoustic pressure: < 0.1 Mpa |
| Ultrasound output power: ≤20mW |
| FHR display range: 50 – 210 bpm |
| Resolution: 1 bpm |
| Precision: ± 2 bpm |
| Curve display range: 90 – 190 bpm |
| Battery: AAx2 |
| Work mode: continuous (The device can work continuously for over 4 hours) |
| Dimension: 142.6mm x 40.5mm x 42.5 mm |
| Weight: 130±5 g |
| Ultrasound coupling agent requirement: density = 1.0g/cm ² ; speed ≤1.7m/s; impedance ≤1.7×10 ⁶ g/cm ² .s; attenuation ≤0.02dB/mm |
| P _r < 1MPa; Iob < 20 mW/cm ² ; Ispta < 100mW/cm ² |

Manufacturing date: See the label.

Device life expectancy: 5 years

Waterproofing grade: IP22

Safety type: Internally powered equipment, type BF applied part

Software version: 1.0

Operation conditions: Temperature: 5°C to 40°C;

Humidity: ≤80%RH; non-condensing

Atmospheric pressure: 70kpa to 106kpa

Transportation & Storage conditions:

Temperature: -20°C to 55°C;

Humidity: 10%RH – 93%RH; non-condensing

Atmospheric pressure: 50kpa to 106kpa; indoor ventilated place that has no corrosive gas

Equipment heating time

-the time required for the equipment to warm from the minimum storage temperature between uses until it is ready for intended use: 30min.

-the time required for the equipment to cool from the maximum storage temperature between uses until it is ready for intended use: 30min.

5.2 Troubleshooting

| Symptom | Possible cause | Troubleshooting |
|------------------|----------------|-----------------------|
| Power-on failure | Low battery | Charge the instrument |

| Symptom | Possible cause | Troubleshooting |
|-----------------------------|--|---|
| No sound | Low volume Low power | Increase the volume Replace the battery |
| Fetal heart cannot be found | Low volume The coupling agent is not coated | Increase the volume Coat the coupling agent or water |
| Low sensitivity | Incorrect probe location The coupling agent is not coated | Adjust the probe location Coat a proper amount of coupling agent |
| Bluetooth connection failed | Bluetooth on phone is not turned on | Manually turn on the Bluetooth function of the mobile phone |

EMC Information

1* WARNING: Use of this equipment adjacent to or stacked with other equipment should be avoided because it could result in improper operation. If such use is necessary, this equipment and the other equipment should be observed to verify that they are operating normally."

2* WARNING: Use of accessories, transducers and cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result in improper operation."

3* WARNING: Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the ME equipment, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result."

Table 1

| declaration - electromagnetic emission | |
|---|----------------|
| Emissions test | Compliance |
| RF emissions CISPR 11 | Group 1 |
| RF emissions CISPR 11 | Class B |
| Harmonic emissions IEC 61000-3-2 | Not applicable |
| Voltage fluctuations/flicker emissions IEC 61000-3-3 | Not applicable |

Table 2

| declaration - electromagnetic immunity | | |
|--|--|--|
| Immunity test | IEC 60601 test level | Compliance level |
| Electrostatic discharge (ESD) IEC 61000-4-2 | ±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air | ±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air |
| Electrical fast transient/burst IEC 61000-4-4 | ± 2 kV for power supply lines ± 1 kV for input/output lines | Not applicable |
| Surge IEC 61000-4-5 | ± 0.5kV, ± 1 kV line(s) to lines ± 0.5kV, ± 1 kV, ± 2 kV line(s) to earth | Not applicable |
| Voltage dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11 | 0 % UT; 0.5 cycle At 0°, 45°, 90°, 135°, 180°, 225°, 270° and 315° 0 % UT; 1 cycle and 70 % UT; 25/30 cycles Single phase: at 0° 0 % UT; 250/300 cycles | Not applicable |
| Power frequency (50/60 Hz) | 30 A/m | 30 A/m |

| | | |
|--|--|--|
| magnetic field IEC 61000-4-8 | | |
| NOTE: UT is the a.c. mains voltage prior to application of the test level. | | |

Table 3

| declaration - electromagnetic immunity | | |
|--|---|------------------|
| Immunity test | IEC 60601 test level | Compliance level |
| Conducted RF IEC 61000-4-6 | 3 V 0.15 MHz to 80 MHz 6 V in ISM bands between 0.15 MHz and 80 MHz | Not applicable |
| Radiated RF IEC 61000-4-3 | 10V/m 80 MHz to 2.7 GHz | 10V/m |

Table 4

| declaration - IMMUNITY to proximity fields from RF wireless communications equipment | | | | |
|--|----------------------------------|-------------------------------|---------------|------------------|
| Immunity test | IEC60601 test level | | | Compliance level |
| | Test frequency | Modulation | Maximum power | |
| Radiated RF IEC 61000-4-3 | 385 MHz | **Pulse Modulation : 18Hz | 1.8W | 27 V/m |
| | 450 MHz | *FM+ 5Hz deviation: 1kHz sine | 2 W | 28 V/m |
| | 710 MHz 745 MHz 780 MHz | **Pulse Modulation : 217Hz | 0.2 W | 9 V/m |
| | 810 MHz 870 MHz 930 MHz | **Pulse Modulation : 18Hz | 2 W | 28 V/m |
| | 1720 MHz 1845 MHz 1970 MHz | **Pulse Modulation : 217Hz | 2 W | 28 V/m |
| | 2450 MHz | **Pulse Modulation : 217Hz | 2 W | 28 V/m |
| | 5240 MHz 5500 MHz 5785 MHz | **Pulse Modulation : 217Hz | 0.2 W | 9 V/m |
| | | | | |

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

Note** - The carrier shall be modulated using a 50 % duty cycle square wave signal.

FCC Statement

This equipment has been tested and found to comply with the limits for a Class B digital device, pursuant to Part 15 of the FCC Rules. These limits are designed to provide reasonable protection against harmful interference in a residential installation. This equipment generates uses and can radiate radio frequency energy and, if not installed and used in accordance with the instructions, may cause harmful interference to radio communications. However, there is no guarantee that interference will not occur in a particular installation. If this equipment does cause harmful interference to radio or television reception, which can be determined by turning the equipment off and on, the user is encouraged to try to correct the interference by one or more of the following measures:

-- Reorient or relocate the receiving antenna.

-- Increase the separation between the equipment and receiver.

-- Connect the equipment into an outlet on a circuit different from that to which the receiver is connected.

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

This device complies with part 15 of the FCC Rules. Operation is subject to the following two conditions:

(1) This device may not cause harmful interference, and (2) this device must accept any interference received, including interference that may cause undesired operation.

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

* RF warning for Portable device:

The device has been evaluated to meet general RF exposure requirement. The device can be used in portable exposure condition without restriction.

IC STATEMENT

This device contains licence-exempt transmitter(s)/receiver(s) that comply with Innovation, Science and Economic Development Canada's licence-exempt RSS(s). Operation is subject to the following two conditions:

(1) This device may not cause interference.

(2) This device must accept any interference, including interference that may cause undesired operation of the device

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authorities of your Member State.

6.3 Manufacturer

Manufacturer: Shenzhen Jumper Medical Equipment Co., Ltd
Address: D Building, No. 71, Xintian Road, Fuyong Street, Baoan, Shenzhen, Guangdong, China, 518103
Tel: +86-755-26696279
E-mail: info@jumper-medical.com
Website: www.jumpermed.com
www.jumper-medical.com

6.4 Authorized European Representative

 MedPath GmbH

Mies-van-der-Rohe-Strasse 8, 80807 Munich, Germany

FCC

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