

# Wrist Blood Pressure Monitor - MODEL KD-735 (ELECTRONIC SPHYGMOMANOMETER)

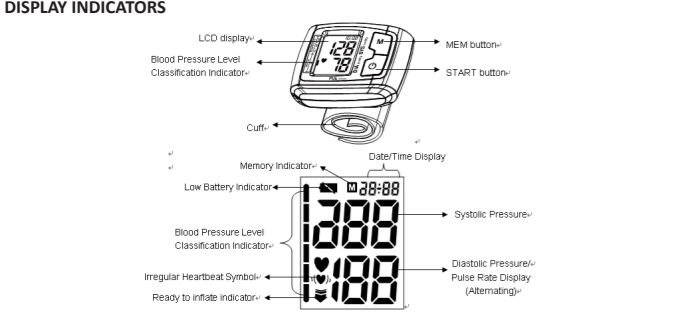
## Instruction Manual

**IMPORTANT INFORMATION**  
Please read this instruction manual carefully before using the product. This manual is for purchasing the Wrist Blood Pressure Monitor. Please retain this Instruction Manual for reference.

**NORMAL BLOOD PRESSURE FLUCTUATION**  
Blood pressure is affected by various factors, including excitement, stress, body position, and physical activities such as eating, drinking, smoking, or even taking a blood pressure measurement. As a result, it is unusual to obtain identical blood pressure readings multiple times. Blood pressure fluctuates constantly throughout the day and night. Typically, it continues to rise during the day and peaks while most people are awake and active. It then drops in the evening, reaching its lowest between midnight and 3 a.m., while most people sleep.

Considering the above information, measuring your blood pressure at approximately the same time every day is recommended. Taking measurements more often than necessary may cause an injury due to blood flow interference, so please always rest at least 60 to 90 seconds between measurements to allow the blood circulation in your arm to recover.

**BOX CONTENTS**  
1 x Blood Pressure Monitor With Attached Wrist Cuff  
1 x Instruction Manual  
1 x Plastic Bag



**INTENDED USE**  
Fully Automatic Electronic Blood Pressure Monitor is for use by medical professionals at or home and is a non-invasive blood pressure measurement system intended to measure the diastolic and systolic blood pressures and pulse rate of an adult individual by using a non-invasive technique in which an inflatable cuff is wrapped around the left wrist. The cuff circumference is limited to 14cm-19.5cm (approx. 5.5"-7.7").

**CONTRAINDICATION**  
This blood pressure monitor (electronic sphygmomanometer) is not suitable for people with severe arrhythmia.

**PRODUCT DESCRIPTION**  
Based on oscillometric methodology and a silicon-integrated pressure sensor, blood pressure and pulse rate can be measured automatically and non-invasively. Users can operate the device themselves. The liquid crystal display (LCD) shows blood pressure and pulse rate. This blood pressure monitoring device can store up to 60 readings for each of two different users, along with the date and time of each measurement. This blood pressure monitor has been designed in accordance with the requirements of ISO 81060-2:2018.

- SPECIFICATIONS**
- 1. Product name: Wrist Blood Pressure Monitor
  - 2. Model: KD-735
  - 3. Classification: Internally powered, Type BF applied part, IP22, No AP or APG, Continuous operation
  - 4. Machine size: Approx. 85mm x 64.5mm x 28mm (3.3" x 2.5" x 1.1")
  - 5. Cuff circumference: 14cm - 19.5cm (5.5" - 7.7")
  - 6. Weight: Approx. 110g (3.9 oz.) (exclude batteries)
  - 7. Measuring method: Oscillometric method
  - 8. Memory volume: Two users, 60 measurements each
  - 9. Power source: batteries: 2 x 1.5V AAA
  - 10. Measurement range: Cuff pressure: 0 to 300 mmHg
- Cuff: 3 years (with 3 uses daily)
- 11. Accuracy: Pressure: ±3 mmHg  
Pulse rate: Less than 60: ±3 bpm  
More than 60 (incl.): ±5%
  - 12. Environmental temperature for operation: 5°C-40°C (41°F-104°F)
  - 13. Environmental humidity for operation: ≤85%RH
  - 14. Environmental temperature for storage and transport: -20°C-55°C (-4°F-131°F)
  - 15. Environmental humidity for storage and transport: ≤90%RH
  - 16. Environmental pressure: 80kPa-105kPa
  - 17. Battery life: Approx 270 times
  - 18. Product life: Monitor: 3 years

**IMPORTANT SAFETY INFORMATION**

- Warning: Indicates a potentially hazardous situation which, if not avoided, could result in death or serious injury.**
- The device should not be used for patients with artificial hearts or lungs. The device should not be used for neonates, infants, children or persons who cannot express themselves. This device has not been validated for use on pregnant patients.
  - The device should not be used for patients with poor peripheral circulation, noticeably low blood pressure, or low body temperature.
  - Consult your physician before using the device for any of the following conditions: common arrhythmias, such as atrial or ventricular premature beats or atrial fibrillation, arterial sclerosis, poor perfusion, diabetes, pre-eclampsia, and renal diseases.
  - Do not use this device in a moving vehicle.
  - Do not use this device if you are allergic to plastic/rubber.
  - Please do not share the cuff to prevent the risk of infection and cross-contamination.
  - Do not use a cuff other than the one supplied by the manufacturer. Disregarding this safety instruction may bring about a biocompatible hazard, result in measurement error.
  - Never let children or persons incapable of expressing themselves independently use the device. Keep the device safely stored and inaccessible to children to prevent them from swallowing the batteries or other small parts.
  - Keep the cuff tube away from children to avoid the risk of strangulation or asphyxiation.
  - As the alternative small-bore connector used for this medical device is designed differently from the connector specified in the ISO 80369 series, the user will need to take steps to reduce the risk of incorrectly connecting.
  - See the section regarding ELECTROMAGNETIC COMPATIBILITY INFORMATION for information regarding potential electromagnetic interference (EMI) or other interference between the device and other devices.

**Caution: Indicates a potentially hazardous situation which, if not avoided, may result in minor or moderate injury to the user.**

- Stay quiet and calm, and rest for five minutes before taking your blood pressure measurement. Relax for a minimum of 1 to 1.5 minutes between measurements to allow the blood circulation in your arm to recover.
- Do not speak or move your body or arm during the measurement. Motion, trembling, and shivering during measurement may affect the result.
- Prolonged overinflation (cuff pressure exceeds 300 mmHg or is above 15 mmHg for longer than 3 minutes) of the bladder may cause ecchymoma (a tumor-like swelling composed of extravasated blood) of the arm.
- The device might not meet its performance specifications or cause safety hazards if stored or used outside the specified temperature and humidity ranges listed in the specifications.
- Consult your physician before use if any of the following scenarios are applicable:
  - 1) The cuff will be applied over a wound or inflammation disease;
  - 2) The cuff will be applied on any arm where intravascular access or therapy, or an arterio-venous (A-V) shunt, is present;
  - 3) The cuff will be applied to the arm on the same side as a mastectomy or lymph node clearance;
  - 4) The device will be used simultaneously with other monitoring medical equipment on the same arm.
- Blood pressure measurements determined by this device are equivalent to those obtained by a trained observer using the cuff/stethoscope auscultation method, within the limits prescribed by the American National Standard Institute, electronic or automated sphygmomanometers.
- A signal will be displayed if the blood pressure measurement procedure detects an irregular heartbeat (IHB) brought on by common arrhythmias.
- Under this condition, the electronic sphygmomanometer can keep functioning, but the results may not be accurate. Please consult your physician for a precise assessment.
- There are conditions under which the signal of IHB will be displayed:
  - 1) The coefficient of variation of pulse period is ≥25 percent.
  - 2) The difference in adjacent pulse periods is ≥ 0.14 seconds, and the number of such pulses is more than 53 percent of the total number.
- Please check the condition of the arm being used to ensure that the device is not impairing the patient's blood circulation when in use.
- All serious accidents concerning the medical device supplied by us must be reported to the manufacturer and competent authority of the member state where your registered office is located.

**SETUP AND OPERATION**

**1. INSTALLING BATTERY**

- Open the battery cover at the back of the device.
- Insert two "AAA" batteries. Make sure the batteries are inserted according to the positive and negative marks ("+" and "-") printed in the battery housing.
- Close the battery cover.

**Note:**  
When LCD shows a low battery symbol, replace all batteries with new ones. Rechargeable batteries are not suitable for this device.

**Warning:** Avoid getting battery fluid in your eyes. If battery fluid gets in your eyes, immediately rinse with plenty of clean water and consult with your physician.

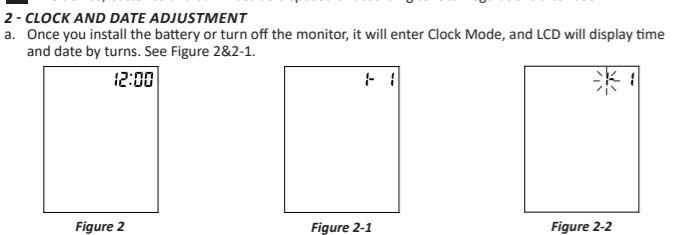
The negative (-) side of the battery should be touching the spring.

Ensure the battery cover is intact and not damaged before installing the battery.

The device, batteries and cuff must be disposed of according to local regulations after use.

**2 - CLOCK AND DATE ADJUSTMENT**

- Once you install the battery or turn off the monitor, it will enter Clock Mode, and LCD will display time and date by turns. See Figure 2&2-1.



While the monitor is in Clock Mode, pressing both the "START" and "MEM" button simultaneously, a beep is heard and the month will blink at first. See Figure 2-2. Press the button "START" repeatedly, the day, hour and minute will blink in turn. While the number is blinking, press the button "MEM" to increase the number. Keep on pressing the button "MEM", the number will increase fast.

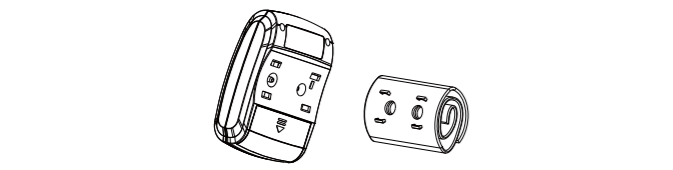
You can turn off the monitor by pressing "START" button when the minute is blinking, then the time and date is confirmed.

The monitor will turn off automatically after 1 minute of no operation, with the time and date unchanged.

Once you change the batteries, you should readjust the time and date.

**3 - CONNECTING THE CUFF TO THE DEVICE**

The cuff is attached to the device when it is packaged. Should the cuff become unattached, align the two plugs and four brackets of the cuff with the plug sockets and bracket sockets of the monitor and press the cuff to the monitor until the plugs and brackets are securely attached.



**4 - APPLYING THE CUFF**

- Place the cuff around a bare left wrist 1-2cm above the wrist joint on the palm side of the wrist.
- While seated, place the arm with the cuffed wrist in front of your body on a desk or table with the palm up. If the cuff is correctly placed, you can read the LCD display.
- The cuff must be neither too tight nor too loose.



**Note:**

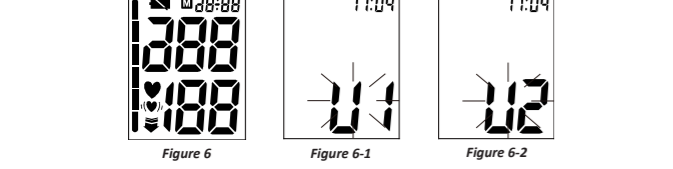
- Please refer to the cuff circumference range in "SPECIFICATIONS" to ensure appropriate usage.
- Always measure on the same arm for consistency.
- Do not apply the cuff if the arm has inflammation, acute diseases, or skin wounds.

**5 - BODY POSTURE DURING MEASUREMENT SITTING DURING MEASUREMENT:**

- Sit with your feet flat on the floor and avoid crossing your legs.
- Extend your arm with your palm facing up, resting comfortably on a flat surface.
- Position the cuff to be at the same level as your heart.

**6 - TAKING YOUR BLOOD PRESSURE READING**

- After applying the cuff and your body is in a comfortable position, press the "START" button. A beep is heard and all display characters are shown for self-test. See Figure 6. Please contact the service center if segment is missing.



Then the current memory bank (U1 or U2) is blinking. See Figure 6-1. Press "MEM" button to change over to other bank. See Figure 6-2. Confirm your selection by pressing "START" button. The current bank can also be confirmed automatically after 5 seconds with no operation.

After selecting the memory bank, the monitor starts to seek zero pressure. See Figure 6-3.

The monitor inflates the cuff until sufficient pressure has built up for a measurement. Then the monitor slowly releases air from the cuff and carries out the measurement. Finally the blood pressure and pulse rate will be calculated and displayed on the LCD screen separately. Irregular heartbeat symbol (if any) will blink. See Figure 6-4&6-5. The result will be automatically stored in the current memory bank.

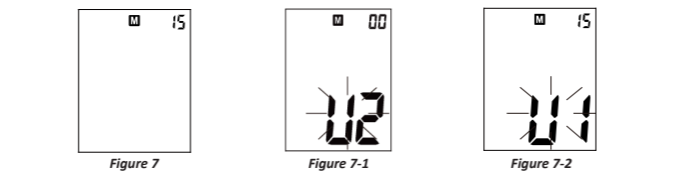


After measurement, the monitor will turn off automatically after 1 minute of no operation. Alternatively, you can press the "START" button to turn off the monitor manually.

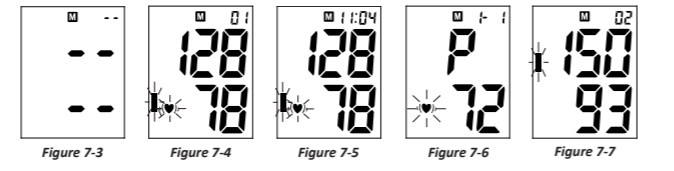
During measurement, you can press the "START" button to turn off the monitor manually. Note: Please consult a health care professional for interpretation of pressure measurements.

**7 - DISPLAYING STORED RESULTS**

- After the measurement, you can review the measurements in the current memory bank by pressing button "MEM". Now the LCD displays the amount of the results in the current bank. See Figure 7.



Alternatively, press "MEM" button in Clock Mode to display the stored results. The current memory bank will blink and the amount of results in this bank will be displayed. See Figure 7-1. Press "START" button to change over to other bank. See Figure 7-2. Confirm your selection by pressing "START" button. The current bank can also be confirmed automatically after 5 seconds with no operation. After selecting the memory bank, the LCD will display the average value of the last three results in this bank. If no result stored, LCD will show dashes as Figure 7-3.

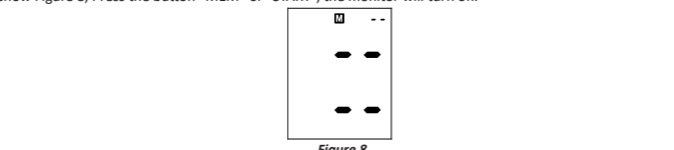


When the average is displayed, press the "MEM" button, the most recent result will be displayed. See Figure 7-4. Followed by, the blood pressure and pulse rate will be shown separately. Irregular heartbeat symbol (if any) will blink. See Figure 7-5&7-6. Press "MEM" button again to review the next result. See Figure 7-7. In this way, repeatedly pressing the "MEM" button displays the respective results measured previously.

When displaying the stored results, the monitor will turn off automatically after 1 minute of no operation. You can also press the button "START" to turn off the monitor manually.

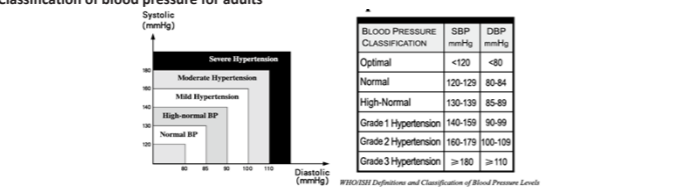
**8 - DELETING MEASUREMENTS FROM THE MEMORY**

When any result (except average reading of the last three results) is displaying, keeping on pressing button "MEM" for three seconds, all results in the current memory bank will be deleted after three "beep". LCD will show Figure 8. Press the button "MEM" or "START", the monitor will turn off.



**9 - ASSESSING HIGH BLOOD PRESSURE FOR ADULTS**

The World Health Organization has established the following guidelines for assessing high blood pressure (regardless of age or gender). Please note that other factors (e.g., diabetes, obesity, smoking, etc.) must be considered. Consult with your physician for an accurate assessment. Only change any existing course of treatment by yourself after first seeking the advice of a medical professional.



The guidelines are not intended to be used as a basis for self-diagnosis or emergency conditions but only to differentiate between general classifications of blood pressure levels.

**TROUBLESHOOTING (1)**

PROBLEM	POSSIBLE CAUSE	SOLUTION
LCD Display shows abnormal result	The cuff position was not correct or it was not properly tightened	Apply the cuff correctly and try again
	Body posture was not correct during testing	Review the "BODY POSTURE DURING MEASUREMENT" sections of the instructions and re-test.
	Speaking, arm or body movement, angry, excited or nervous during testing	Re-test when calm and without speaking or moving during the test
	Irregular heartbeat (arrhythmia)	It is inappropriate for people with serious arrhythmia to use this Electronic Sphygmomanometer.

**TROUBLESHOOTING (2)**

PROBLEM	POSSIBLE CAUSE	SOLUTION
LCD shows low battery symbol	Low Battery	Change the batteries
LCD shows "Er 0"	Pressure system is unstable before measurement	Don't move and try again.
LCD shows "Er 1"	Fail to detect systolic pressure	
LCD shows "Er 2"	Fail to detect diastolic pressure	
LCD shows "Er 3"	Pneumatic system blocked or cuff is too tight during inflation	Apply the cuff correctly and try again
LCD shows "Er 4"	Pneumatic system leakage or cuff is too loose during inflation	
LCD shows "Er 5"	Cuff pressure above 300mmHg	
LCD shows "Er 6"	More than 3 minutes with cuff pressure above 15 mmHg	Measure again after five minutes. If the monitor is still abnormal, please contact the local distributor or the factory.
LCD shows "Er 7"	EEPROM accessing error	
LCD shows "Er 8"	Device parameter checking error	
LCD shows "Er A"	Pressure sensor parameter error	
No response when you press button or load battery.	Incorrect operation or strong electromagnetic interference.	Take out batteries for five minutes, and then reinstall all batteries.

- MAINTENANCE**
- Avoid dropping or subjecting the device to substantial impacts.
  - Avoid high temperatures and prolonged exposure to direct sunlight. Do not immerse the device in water, which will damage it.
  - Changes or modifications not approved by the manufacturer will void the user's warranty. Do not disassemble or attempt to repair the device or its components.
  - Remove the batteries if the device is not used for longer than one month to avoid damage due to battery leakage.
  - It is recommended that the device's performance be checked every two years.
  - Clean the device with a dry, soft cloth or a soft cloth dampened with water, disinfectant alcohol, or diluted detergent (wringing out the cloth to remove as much liquid as possible before wiping the device).
  - Please keep the cuff clean. If the cuff becomes dirty, remove it from the device and clean it by hand with mild detergent, then rinse it thoroughly with cold water. Never dry the cuff in a clothes dryer or iron it. For personal use, it is recommended that the cuff be cleaned after it has been used approximately 200 times. Disinfecting is recommended if the cuff is used in a hospital or a clinic. Wipe the cuff's inner side (the side that contacts the skin) with a soft cloth, lightly moistened with Ethyl alcohol (75 to 90 percent). Then, air-dry the cuff.
  - We can provide product circuit diagrams and repairable component information to qualified maintenance service personnel if necessary.
  - Please wait when moving the device between extreme temperatures (e.g., storage, during transport) to a normal operating environment. The device takes approximately two hours to warm up or cool down before use.
  - The device shall not be serviced or maintained while in use.

**ELECTROMAGNETIC COMPATIBILITY INFORMATION**

- This medical device meets the following essential performance requirements:
  - Limits of the error of the cuff pressure indication.
  - Reproducibility of the blood pressure determination.
- When EMI affects the above performance, please stop using the device.
- 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 usually operate.
- 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.
- Equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the device, including cables specified by the manufacturer. Otherwise, degradation of the performance of this equipment could result."

**Table 1 - Emissions**

Phenomenon	Compliance	Electromagnetic Environment
Radiated RF emissions	CISPR 11 User 1, Class B	Home healthcare environment
Harmonic distortion	IEC 61000-3-2	This device is powered by batteries
Voltage fluctuations and flicker	IEC 61000-3-3 NA	This device is powered by batteries

**Table 2 - Enclosure Port**

Phenomenon	Basic EMC Standard	Immunity Test Levels
		Home Healthcare Environment
Electrostatic Discharge	IEC 61000-4-2	±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air
Radiated RF EM field	IEC 61000-4-3	10 V/m 80 MHz to 2.7 GHz 80% AM at 1 kHz
Proximity fields from RF wireless communications equipment	IEC 61000-4-3	Refer to Table 3
Rated power frequency magnetic fields	IEC 61000-4-8	30 A/m 50 Hz or 60 Hz

**Table 3 - Proximity Fields from RF Wireless Communications Equipment**

Test frequency (MHz)	Band (MHz)	Immunity Test Levels
		Professional healthcare facility environment
385	380-390	Pulse modulation 18 Hz, 27 V/m
450	430-470	FM, ±5 kHz deviation, 1 kHz sine, 28 V/m
710		
745	704-787	Pulse modulation 217 Hz, 9 V/m
780		
810		
870	800-960	Pulse modulation 18 Hz, 28 V/m
930		
1720		
1845	1700-1990	Pulse modulation 217 Hz, 28 V/m
1970		
2450	2400-2570	Pulse modulation 217 Hz, 28 V/m
5240		
5500	5100-5800	Pulse modulation 217 Hz, 9 V/m
5785		

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

**GIMA WARRANTY TERMS**  
The Gima 12-month standard 828 warranty applies

Date of issue: Apr. 10, 2025

Indice dei simboli - Symbol index - Index des symboles - Índice de símbolos - Índice de símbolo - Index de simbol - Ευρετήριο συμβόλων - Symbolindex - Index de simbol - Szimbólum index - زيورلا برهف	
	IT - Data di fabbricazione GB - Date of manufacture FR - Date de fabrication ES - Fecha de fabricación PT - Data de fabrico DE - Herstellungsdatum GR - Ημερομηνία παραγωγής PL - Data produkcyj SE - Tillverkningsdatum RO - Data fabricației HU - Gyártás dátuma AR - تاريخ التصنيع
	IT - Fabricante GB - Manufacturer FR - Fabricant ES - Fabricante PT - Fabricante DE - Hersteller GR - Παράγωγός PL - Producent SE - Tillverkare RO - Producător HU - Gyártó AR - الشركة المصنعة
	IT - Conservare al riparo dalla luce solare GB - Keep away from sunlight FR - À conserver à l'abri de la lumière du soleil ES - Conservar al amparo de la luz solar PT - Guardar ao abrigo da luz solar DE - Vor Sonneneinstrahlung geschützt lagern GR - Κρατήστε το μακριά από ηλιακή ακτινοβολία PL - Przechowywać z dala od światła słonecznego SE - Skyddas från solljus RO - A se păstra ferit de razele soarelui HU - Napfénytől védve tárolandó AR - يحفظ بعيداً عن ضوء الشمس
	IT - Importato da GB - Imported by FR - Importé par ES - Importado por PT - Importado por DE - Eingeführt von GR - Εισαγύγνται από PL - Importowane przez SE - Importerad av RO - Importat de HU - Importálta AR - مستورد عن طريق
	IT - Conservare in luogo fresco ed asciutto GB - Keep in a cool, dry place FR - À conserver dans un endroit frais et sec ES - Conservar en un lugar fresco y seco PT - Armazenar em local fresco e seco DE - An einem kühlen und trockenen Ort lagern GR - Διατηρείται σε όψωρο και στεγνό μέρος PL - Przechowywać w suchym miejscu SE - Förvara på svalt och torrt ställe RO - A se păstra într-un loc răcoșos și uscat HU - Száraz, hűvös helyen tárolandó AR - يحفظ بعيداً عن ضوء الشمس
	IT - Attenzione: Leggere e seguire attentamente le istruzioni (avvertenze) per l'uso GB - Caution: read instructions (warnings) carefully FR - Attention: lisez attentivement les instructions (avertissements) ES - Precaución: lea las instrucciones (advertencias) cuidadosamente PT - Cuidado: leia as instruções (avisos) cuidadosamente DE - Achtung: Anweisungen (Warnungen) sorgfältig lesen GR - Προσοχή: διαβάστε προσεκτικά τις οδηγίες (επιδείξεις) PL - Ostrzeżenie - Zobacz instrukcję obsługi SE - Varsamhet: läs anvisningarna (varningar) noga RO - Atenție: Citiți și respectați cu atenție instrucțiunile (avertismentele) de utilizare HU - Figyelem: Figyelmesen olvassa el és kövesse a használati utasításokat (figyelmeztetéseket) AR - ليس معقم
	IT - Seguire le istruzioni per l'uso GB - Follow instructions for use FR - Suivez les instructions d'utilisation ES - Siga las instrucciones de uso PT - Siga as instruções de uso DE - Folgen Sie den Anweisungen GR - Ακολουθήστε τις οδηγίες χρήσης PL - Patrz podaćcznik użytkownika SE - Följ bruksanvisningen RO - Respectați instrucțiunile de utilizare HU - Kövesse a használati utasításokat AR - اتبع التعليمات للاستخدام
	IT - Limite di umidità GB - Humidity limit FR - Limite d'humidité ES - Limite de humedad PT - Limite de humidade DE - Feuchtigkeitsgrenzwert GR - Όριο υγρασίας PL - Granica wilgotności SE - Fuktighetsgräns RO - Limită de umiditate HU - Páratartalom határérték AR - حد نسبة الرطوبة
	IT - Limite di temperatura GB - Temperature limit FR - Limite de température ES - Limite de temperatura PT - Limite de temperatura DE - Temperaturgrenzwert GR - Διατηρείται μεταξύ -10 και 49°C PL - Przechowywać pomiędzy -10 °C SE - Lagras mellan och °C RO - A se păstra la temperaturi cuprinse între și °C HU - és °C között tárolandó AR - يحفظ بين و درجة مئوية
	IT - Numero di lotto GB - Lot number FR - Numéro de lot ES - Número de lote PT - Número de lote DE - Chargennummer GR - Αριθμός παρτίδας PL - Kod partii SE - Satsnummer RO - Număr de lot HU - Tételszám AR - رقم الدفعة
	IT - Codice prodotto GB - Product code FR - Code produit ES - Código productu PT - Código produto DE - Erzeugniscode GR - Κωδικός προϊόντος PL - Numer katalogowy SE - Produktkod RO - Cod produs HU - Termékkód AR - كود المنتج
	IT - Smaltimento RAEE GB - WEEE disposal FR - Disposition DEEE ES - Disposición WEEE PT - Disposição REEE DE - Beseitigung WEEE GR - Διάθεση WEEE PL - Oddzielna zbiórka dla tego urządzenia SE - Avfallshantering av elektrisk och elektronisk utrustning (WEEE) RO - Eliminare DEEE HU - RAEE szerinti ártalmatlanítás AR - التخلص WEEE
	IT - Dispositivo medico conforme al regolamento (UE) 2017/745 GB - Medical Device compliant with Regulation (EU) 2017/745 FR - Dispositif médical conforme au règlement (UE) 2017/745 ES - Producto sanitario conforme con el reglamento (UE) 2017/745 PT - Dispositivo médico em conformidade com o regulamento (UE) 2017/745 DE - Medizinprodukt im Sinne der Verordnung (EU) 2017/745 GR - Ιατρική συσκευή σύμφωνα με την ΚΑΝΟΝΙΣΜΟΣ (ΕΕ) 2017/745 PL - Wyrób medyczny zgodny z Rozporządzeniem (UE) 2017/745 SE - Den medicintekniska produkten överensstämmer med förordning 2017/745 (EU) RO - Dispozitiv medical conform regulamentului (UE) 2017/745 HU - A 2017/745/EU rendelethez megfelelő orvostechnikai eszköz AR - جهاز طبي يتوافق مع التوجيه (UE) 2017/745
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	IT - Rappresentante autorizzato GB - Authorized representative in the European Union FR - Représentant autorisé ES - Representante autorizado PT - Representante autorizado DE - Autorisierter Vertreter GR - εξουσιοδοτημένος αντιπρόσωπος PL - Upoważniony przedstawiciel SE - Auktoriserad representant HU - Meghatalmazott képviselő RO - Reprezentant autorizat AR - الممثل المعتمد
	IT - Parte applicata di tipo BF GB - Type BF applied part FR - Appareil de type BF ES - Aparato de tipo BF PT - Aparelho de tipo BF DE - Gerätetyp BF GR - Συσκευή τύπου BF PL - Z części typu BF SE - Typ BF tillämpad del HU - Meghatalmazott képviselő RO - Représentant autorizat AR - جهاز من النوع BF
	IT - Identificatore univoco del dispositivo GB - Unique device identifier FR - Identifiant unique de l'appareil ES - Identificador de dispositivo único PT - Identificador exclusivo do dispositivo DE - Unique Device Identifier (Eindeutige Kennung des Geräts) GR - Μοναδικό αναγνωριστικό συσκευής PL - Unikalny identyfikator urządzenia SE - Unik identifierare för enheten RO - Identificatorul unic al dispozitivului HU - Az eszköz egyedi azonosítója AR - معرف فريد للجهاز
	IT - Grado di protezione dell'involucro GB - Covering Protection rate FR - Degré de protection de l'enveloppe ES - Tasa de protección de cobertura PT - Grau de protecção do invólucro DE - Deckungsschutzrate GR - Δείκτης στεγανότητας PL - Stopień ochrony obudowy SE - Skyddsgrad RO - Grad de protecție asigurată prin carcasă HU - A csomagolás védelmi szintje AR - مؤشر الغاذية
	IT - Numero di serie GB - Serial number FR - Numéro de série ES - Número de serie PT - Número de série DE - Seriennummer GR - Σειριακός αριθμός PL - Numer serijny SE - Serienummer RO - Număr de serie HU - Sorozatszám AR - الرقم التسلسلي
	IT - Non sicuro in ambiente RM GB - MR Unsafe FR - IRM dangereuse ES - MR inseguro PT - Não seguro para RM DE - "MR Unsicher" (nicht für den Einsatz im MR geeignet) GR - Μη ασφαλές για MR PL - Niebezpieczne w środowisku RM SE - Inte säker i MR-miljö RO - Nesigur pentru RM HU - MR nem biztonságos AR - غير آمن للرنين المغناطيسي



- SMART-WRIST**
- MISURATORE DI PRESSIONE AUTOMATICO DA POLSO
  - WRIST AUTOMATIC BLOOD PRESSURE MONITOR
  - TENSIOÛTRE AUTOMATIQUE AU POIGNET
  - MONITOR AUTOMÁTICO DE PRESIÓN ARTERIAL DE MUÑECA
  - MONITOR AUTOMÁTICO DE PRESSÃO ARTERIAL DE PULSO
  - AUTOMATISCHES HANDGELENK-BLUTDRUCKMESSGERÄT
  - AUTOMATYCZNY CIŚNIENIOMIERNY NADGARSTKOWY
  - ΑΥΤΟΜΑΤΟ ΠΙΕΣΟΜΕΤΡΟ ΚΑΡΠΟΥ
  - AUTOMATISK BLODTRYCKSMÄTARE FÖR HANDELEDEN
  - TENSIOÛTRU AUTOMAT PENTRU ÎNCHEIETURA MĂINII
  - INTELLIGENS AUTOMATA VÉRNYOMÁSMÉRŐ
- جهاز قياس ضغط الدم التلقائي من المعصم

**ATTENTION:** The operators must carefully read and completely understand the present manual before using the product.  
**ATTENZIONE:** Gli operatori devono leggere e capire completamente questo manuale prima di utilizzare il prodotto.  
**AVIS:** Les opérateurs doivent lire et bien comprendre ce manuel avant d'utiliser le produit.  
**ATENCIÓN:** Los operadores tienen que leer y entender completamente este manual antes de utilizar el producto.  
**CUIDADO:** Os operadores devem ler e compreender este manual completamente antes de usar o produto.  
**VORSICHT:** Bediener müssen dieses Handbuch vollständig lesen und verstehen, bevor sie das Produkt verwenden.  
**UWAGA:** Operatorzy muszą w całości przeczytać i zrozumieć niniejszą instrukcję przed użyciem produktu.  
**ΠΡΟΣΟΧΗ:** Οι χειριστές πρέπει να διαβάσουν και να κατανοήσουν πλήρως αυτό το εγχειρίδιο πριν από τη χρήση του προϊόντος.  
**ATTENZIONE:** Operatorii trebuie să citească și să înțeleagă complet acest manual înainte de a utiliza produsul.  
**VIGYÁZAT:** A kezelőknek el kell olvasniuk és meg kell érteniük ezt a kézikönyvet a termék használatá előt.  
**FÖRSIKTIGHET:** Operatörerna måste läsa och förstå denna manual helt innan de använder produkten.  
**تنبيه:** يجب على المتغلين قراءة هذا الدليل وفهمه بالكامل قبل استخدام المنتج.

**GIMA 32918**

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5°C 0%

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M32918-B-M-Rev.03-05-25