OPERATOR'S MANUAL

HeartOn A15

Automated External Defibrillator

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Directive

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- The contents of this manual are subject to change without notice.
- The contents of this manual should be correct. If, for some reason, there are any questionable points, please do not hesitate to contact our service center.
- The manual will be replaced if any pages are missing or collation is incorrect.

Warranty

- Device failure or damage related to the following situations during the guarantee period is not covered by this warranty:
 - Installation, transfer installation, maintenance and repairs by any person other than an authorized Mediana employee or technician specified by Mediana.
 - Damage sustained to the Mediana product(s) caused by product(s) from another company excluding products delivered by Mediana.
 - Damage caused by mishandling and/or misuse is the responsibility of the user.
 - Maintenance and repairs utilizing maintenance components that are not specified by Mediana.
 - Device modifications or use of accessories not recommended by Mediana.
 - Damage caused by accidents or natural disasters (earthquakes, flooding, etc.).
 - Damage resulting from usage where caution statements and operating instructions shown in this manual have not been followed.
 - Damage due to neglect of specified maintenance checks.
- This warranty only covers the hardware of the HeartOn A15. The warranty does not cover the following selections:
 - Whatever damage or loss results from the attachment of accessories or their operation.
 - In the event of a defect in the product, contact our sales outlet or EU representative as noted on the back cover.
- The HeartOn A15 conforms to the EMC standard IEC60601-1-2.

Note: It is possible that using in the vicinity of mobile phone may result in disruption in the AED operation.

Revision History

The documentation part number and revision number indicate its current edition. The revision number changes when a new edition is printed in accordance with the revision history of the documentation. Minor corrections and updates which are incorporated at reprint do not cause the revision number to change. The document part number changes when extensive technical changes are incorporated.

Trademark

Product brand names shown in this manual are likely to be the trademark or registered trademark of the company concerned.

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SAFETY INFORMATION

General Safety Information

This section contains important safety information related to general use of the HeartOn A15. Other important safety information appears throughout the manual. The HeartOn A15 will be referred to as the AED throughout this manual.

Before use, carefully read operator's manual, accessory directions for use, all precautionary information and specifications.

Warning

Warnings alert you to potential serious outcomes (death, injury, or adverse events) to the patient or user.

<u></u> MARNING	As a user of an AED it is essential that you inform Mediana of any incident where your AED is suspected to have caused a death, serious injury or illness. If you have any suspicions that this is the case inform Mediana directly or through your authorized Mediana dealer.
⚠ WARNING	The AED has the capability to deliver therapeutic electrical shocks. The shock can cause serious harm to either operators or bystanders. Caution must be taken to ensure that neither the operators nor bystanders touch the patient when a shock is to be delivered.
⚠ WARNING	The AED has not been evaluated or approved for use in hazardous locations as defined in the National Electrical Code (Articles 500-503). In accordance with the IEC/EN 60601-1 Classifications, the AED is not to be used in the presence of flammable substance/air mixtures.
⚠ WARNING	The AED has been designed to work on unresponsive, non-breathing and pulseless* patients. If the patient is conscious or breathing and regain a pulse, do not use the AED to provide treatment. (*checking pulse corresponds to health care provider)
⚠ WARNING	Touching the patient during the analysis phase of treatment can cause interference with the diagnostic process. Avoid contact with the patient and keep the patient as motionless as possible while ECG analysis is being carried out. The AED will instruct you when it is safe to touch the patient.
⚠ WARNING	Always stand clear of patient when delivering treatment. Defibrillation energy delivered to the patient may be conducted through the patient's body and cause a lethal chock to those touching the patient.
⚠ WARNING	It has been determined that the AED is safe to use in conjunction with oxygen mask delivery systems. However, due to the danger of explosion it is strongly advised that the AED should not be used in the vicinity of explosive gases. This includes flammable anesthetics, concentrated oxygen and gasoline.
⚠ WARNING	The identical pad is used for both Adult and Pediatric (Infant-Child). The Adult mode must be used on patients older than 8 years old. The Pediatric (Infant-Child) mode must be used on patients between 1 and 8 years or less than 25 kg (55lb). Do not use the AED on patient younger than 1 year old.
⚠ WARNING	Proper placement of the pads is critical. Strict observance of pad positioning instructions, as indicated on the labelling is essential. Care

	must be taken to ensure pads are adhered to the patients' skin properly. Air pockets between the adhesive pad and skin must be eliminated. Failure in pad adhesion may hinder effectiveness of therapy or cause excessive skin burns to the patient if a shock is applied. Reddening of the skin may appear after use, this is normal.
⚠ WARNING	The battery of AED is not rechargeable. Do not try to recharge, open, crush, or burn the battery, or it may explode or catch fire.
⚠ WARNING	Do not allow the pads to contact other electrodes or metal parts that are in contact with the patient. Such contact can cause patient skin burns during defibrillation and may divert defibrillating current away from the heart.
⚠ WARNING	Pay attention to possibility of contact with conduction part of electrode, lead line, cable connector, other patient installation part for patient safety.
⚠ WARNING	Do not use this AED near or within puddles of water.
⚠ WARNING	Do not reuse electrodes to many patients.
⚠ WARNING	Use the AED or accessories only as described in this manual. Improper use of the AED can cause death or injury.
⚠ WARNING	Do not use or place the AED in service if the status indicator of AED displays "X".
⚠ WARNING	Keep batteries dry and away from any heat sources (including direct sunlight). If you see any damage or leakage, do not allow the liquid to come in contact with your skin or eyes. If contact has been made, wash the affected area with plenty of water and seek medical advice immediately.
⚠ WARNING	The AED contains an automatic disarm of the stored energy. If the operator has not delivered the energy to a patient, an internal timer will disarm the stored energy. This stored electrical energy can potentially cause death or injury if discharged improperly. Follow all instructions in this manual.

Cautions

Caution statements identify conditions or practices that could result in damage to the equipment or other property.

⚠ CAUTION	The AED may not operate properly if it is operated or stored at conditions outside the ranges stated in this manual.
⚠ CAUTION	The AED was designed to be sturdy and reliable for many different use conditions. However, handling the AED too roughly can damage it or its accessories and will invalidate the warranty. Check the AED and accessories regularly for damage, according to directions.
⚠ CAUTION	Before delivering a shock, it is important to disconnect the patient from non-defibrillation protected electronic devices, such as blood-flow meters, that may not incorporate defibrillation protection. In addition, make sure the pads are not in contact with metal objects such as a bed frame or stretcher.
⚠ CAUTION	The pads pouch shall not be opened until immediately prior to use.
⚠ CAUTION	Do not use or place the AED in service until you have read the AED Operator's manual.
⚠ CAUTION	Do not use or stack the AED with other equipment. If the AED is used or stacked with other equipment, verify proper operation prior to use.
⚠ CAUTION	Handling or transporting the patient during ECG analysis can cause incorrect or delayed diagnosis. If the AED gives a SHOCK ADVISED prompt during such handling or transport, stop the vehicle and keep the patient as still as possible for at least 15 seconds before pressing the Shock button, to allow the AED to reconfirm the rhythm analysis.
⚠ CAUTION	Periodic checks of this AED must be undertaken to ensure among other things that the AED is not damaged in any way.
⚠ CAUTION	The pads are a single use item and must be replaced after each use or if pouch that seals pads has been broken/compromised in any way. If damage is suspected the pads must be replaced immediately.
⚠ CAUTION	Do not use training pads with this AED.
⚠ CAUTION	Carefully observe pacemaker patients. Patient history and physical examination are important in determining the presence of implanted pacemaker. Patient pacemakers may reduce the sensitivity of the AED analysis and errors in detecting shockable rhythms.
⚠ CAUTION	If the pads are attached to the chest firmly, the AED can analyze the exact ECG and prevent the skin burns. But if the pads are overlapped on the patient chest, the pads will not deliver defibrillation energy properly.

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INTRODUCTION

Mediana provides you with a fully configurable AED system to allow you to comply with your chosen SCA treatment protocol. Our current AED is configured to be compliant with the 2021 version of the ERC guidelines on Cardiopulmonary Resuscitation (CPR) and Emergency Cardiovascular Care (ECC). It is recommended to be trained in the appropriate version of the ERC guidelines and the use of your AED configuration. Contact Mediana or your authorized Mediana distributor for further information.

Intended Use for the AED

The AED is intended to be used to treat someone who is unresponsive, non-breathing and pulseless for the adult and pediatric (infant-child) in all area of a hospital, pre-hospital, public access, alternate care and home healthcare environment. AED is designed to easy to use.

Note: The intended patient populations are adult and pediatric (infant-child) (between 1 and 8 years or less than 25 kg (55lb)) can be treated with the appropriate pads.

Note: If you have concerns about your health or an existing medical condition, talk to your doctor. A defibrillator is not a replacement for seeking medical care.

Where can it be used?

The intended environment to use the AED includes home healthcare, public space and hospital. The public space is a social space that is generally open and accessible to people. Roads (including the pavement), public squares, parks, subway station, government buildings, beaches, public libraries, privately owned buildings or property opened to public/visible from sidewalks and any shared spaces of automobiles and other vehicles are typically considered public space. Hospital use typically includes areas such as general care floors, operating rooms, special procedure areas, intensive and critical care areas within the hospital. Hospital-type facilities include physician office-based facilities, sleep labs, skilled nursing facilities, surgical centers, and sub-acute care centers.

Who can use it?

You cannot use the AED to treat yourself. The AED talks the user through each step of treating someone who is in SCA. Responding to SCA may require the user to kneel.

Local Requirements

Check with your state health department to see if there are any local or state requirements about owning and using an AED. You can find contact local supplier or Mediana technical support for learning about your country or state.

Device Tracking

This AED may be subject to tracking requirements by the manufacturer and distributors per local regulation. If there are tracking requirements in your local, please notify your local distributor when the AED has been sold, donated, lost, stolen, exported, or destroyed.

About This Manual

This manual explains how to set up and use the AED.

Read the entire manual including the *Safety Information* section, before you operate the AED.

Identifying the AED Configurations

The following table identifies the AED configurations and how they are indicated. The Reference number and serial number are located on the bottom of the AED.

Configuration	Reference No.	Description
HeartOn A15	A15M-G8-0(E)	AED Standard (8 Action Icons)
HeartOn A15-G4	A15M-G4-0(E)	AED Standard (4 Action Icons)

Note: The alphabet "E" can be added as the last digit of reference number in accordance with the region.

Features for the AED

Physical/Mechanical

The AED is an automated external defibrillator (AED) used for the fast delivery of defibrillation electric shock therapy which can be battery-operated.

Electrical

The AED has a battery which is detachable and non-rechargeable.

Display

The indication is LED indicator that flashes red LED under the relevant action icon.

Auxiliary Input/Output(s)

The AED provides Infrared communication port, SD card ports.

DESCRIPTION OF THE AED

Top and Right Panel Components 1 2 3 8

Figure 1. HeartOn A15: Top and Right Panel Components

10

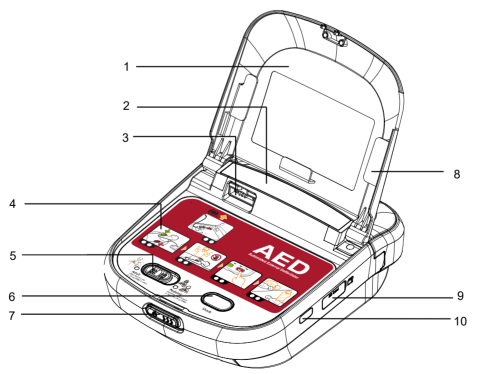
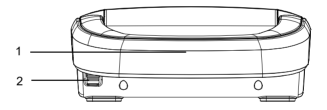


Figure 2. HeartOn A15-G4: Top and Right Panel Components

		Table 1. Top and Right Panel Components			
1	Cover	Cover is used to protect the action icon, the patient mode switch			
		the shock button.			
2	Status indicator	Status indicator displays the AED status, the temperature status			
	Status IIIuicatoi	and the battery status.			
3	Pad connector Pad connector links the pads.				
4	Action icon LED indicator flashes red LED under the relevant action icon.				
		After user distinguish the patient according to patient type, select			
5	Patient mode switch	the patient mode between adult and pediatric (infant-child)			
		patient mode by pushing the patient mode switch.			
		When preparation for electric shock is completed, the shock			
6	Shock button	button will flash. Push the Shock button and then the AED			
		delivers the shock.			
_	0	Slide button is used to open the cover and turn on by pushing			
7	Slide button	the slide button to right.			
8	Pads holder Pads holder is used to fix pads.				
9	SD card port	SD card is used to save the data and update the AED software.			
40	Infrared	Infrared communication port is used to communicate with the			
10	communication port	PC.			

Rear Panel Components



- 1. Handle/Battery
- 2. Battery detachable button

Figure 3. Rear Panel Components

Symbols and Labels

The following symbols may be used in this manual, related documentation, or appear on system components or packaging.

Table 2. Panel and Label Symbols

Symbols	Description	Symbols	Description
0	Ready to use	2	Do not reuse
	Not ready to use	€ 2797	CE mark
	Battery status	0ft 0m	Environmental shipping/storage atmospheric pressure limitations
1	Temperature status	5%	Environmental shipping/storage humidity limitations
Latex Free	Contains no latex	-20°C	Environmental shipping/storage temperature limitations
Σ	Use by date	Ţ	Fragile-handle with care
	Follow instructions for use	<u>†</u>	This way up
***	Manufacturer	Ť	Keep dry
٧	Date of manufacture	1	Type CF – Defibrillator proof
REF	Reference number	IP54	Dust and water resistance
SN	Serial number		Disposal instructions

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SETTING UP THE AED

To ensure accurate performance and prevent AED failure, do not expose the AED to extreme moisture, including direct exposure to rain. Such exposure may cause inaccurate performance or AED failure. Refer to Specification section.

MARNING

Using damaged or expired AED or accessories may cause the AED to perform improperly, and/or injury the patient or the user.

Unpacking and Inspection

The AED is shipped in one carton. Examine the AED including the accessories carefully for evidence of damage. Do not use damaged equipment. Refer to the Maintenance section for instructions on returning damaged items.

Note: Inspect the packaging of accessories to ensure integrity of seals and validity of use by date.

List of Components

The following items are accessories in the package. Optional accessories may be ordered if needed. Contact qualified service personnel or your local supplier for pricing and ordering information.

Table 3. Accessories

Standard Accessories	Qty
HeartOn A15	1
HeartOn A15-G4	
Operator's manual	1
Adult/Pediatric (Infant-Child) Pads (1.8m)	1
Non-rechargeable LiMnO ₂ Battery (15V, 4200mAh)	1
Soft Carry Case	1
SD card	1
Optional Accessories	Qty
HeartOn AED Event Review Software	-
HeartOn AED Event Review Software - User Guide	-
Infrared communication adaptor	-
Mini USB cable	-
Recommended Accessories	Qty
Scissors – for cutting the victim's clothed if needed	-
Disposable gloves – to protect the user	-
A disposable razor- to shave the chest if hair prevents good pads contact	-
A pocket mask or face shield – to protect the user	-
A towel or absorbent wipes - to dry the victim's skin for good pads contact	-

Soft Carry Case

The soft carry case has been designed to allow the AED not to move in the soft carry case by using the AED own handle. The user can check the status indicator of AED without having to open the carry case. The paper with contact information of the nearest emergency medical services can be inserted to the clear cover. The soft carry case has the pocket on the rear side of the carry case for the manual and spare pads. Other two pockets on the right and left side of the carry case are used for spare battery.

SD card

The SD card is inserted into the SD card port on the AED's right panel as described below. The SD card is used to record the history of the AED performance and to update the AED firmware. The recorded history in the SD card can be checked by the HeartOn AED Event Review Software. If you want to use the SD card to use the HeartOn AED Event Review Software or to update the AED firmware, please contact qualified service personnel or your local supplier.

- 1. When the AED is turned on, turn off the AED by closing the cover.
- 2. Open the SD card port cover.
- 3. Insert the SD card into the SD card port.
- 4. Close the SD card port cover.
- 5. If necessary to update the AED through SD card, turn on the AED by pushing **Slide button** to the right.
- 6. After complete the update, automatically turn off itself. Close the cover again.

Event Data

The event data are stored in the SD card. The event data can be read by HeartOn AED Event Review Software.

Note: When the AED does not have an SD card in it or your SD card is unreadable, corrupted, damaged or has some error, the event data are stored in the internal memory as the LED indicator and buzzer sound will go on.

 In case the SD card has a problem while inserting the SD card into the AED during operation, buzzer will sound 4 times. {Repeat twice}

AED records up to 60 minutes of event data in internal memory. The event data do not store any more after 60 minutes data. When more than 50 minutes data are stored, the LED indicator and buzzer sound will go on before hearing the 'Unit ok' voice prompt.

- In case the 50 minutes event data are stored in internal memory during operation, buzzer will sound 5 times. {Repeat twice}
- In case the 50 minutes event data are stored in internal memory during boot up, buzzer will sound 5 times {Repeat twice} and 'action icon – step 5' (HeartOn A15) / 'action icon – step 4' (HeartOn A15-G4) LED indicator will flash.

The event data which are stored in internal memory can be viewed after downloading via SD card in accordance with the following procedure.

- 1. Run the Notepad in Windows. The Notepad window appears with a blank document open.
- 2. Save this empty file in Notepad and name it 'Import Internal Data.txt'.
- 3. Open the SD card on PC.
- 4. In the SD card, create the directory folder and name it 'Update'.
- 5. Copy the carried out 'Import Internal Data.txt' file into the 'Update' directory folder.
- 6. Insert the SD card to SD card port which is located at right side of the AED.
- 7. When the AED is turned on, the event data are downloaded to SD card automatically.
- 8. The downloaded event data can be viewed via the HeartOn AED Event Review Software.

Note: When downloading the stored event data in internal memory to SD card, the event data which are stored in internal memory will be deleted.

Note: If the SD card has some error, the AED can recognize that the SD card is not inserted.

Note: Event data in SD card could be used for further clinical assessment. Please make sure that event data are securely saved in any of storage format when event data is accessed by the HeartOn AED Event Review Software or is uploaded to PC.

Note: It is strongly recommended that the event data in SD card should be uploaded to PC and should be reset before the event data reach 200 events in SD card by periodically checking the HeartOn AED Event Review Software, in order to avoid losing any of event data because the device is designed to stop saving event data when it reaches 200 events in SD card and there is no indication in the device of reaching 200 events in SD card.

Note: If you require any additional information, please refer to HeartOn AED Event Software User guide.

Infrared communication port

Infrared communication port provides wireless communications from the AED to a PC through the Infrared communication data download cable and IR communication adaptor which is connected to PC. The Infrared communication is used to transfer information and to connect to service mode. If you want to use Infrared communication port, please contact qualified service personnel or your local supplier.

Setting up the AED

⚠ WARNING	Use only Mediana-approved and specified parts, accessories, optional parts, consumables, and components. Use of unauthorized accessories may cause the AED to operate improperly and provide false measurements. Follow all labeling instructions on the defibrillation pads and the battery.
⚠ WARNING	Always follow your facility's infection control procedures and applicable regulations when disposing of anything that has been used on patients.
⚠ CAUTION	Do not open the pads from packaging previously until the time of emergency use when pads are used for patient.

Temperature status

Temperature status displays the following description.

- If the self-test is implemented in out-of-range for environmental operation condition above 5 times, the status indicator 'X' will be displayed.
- When the AED with displaying status indicator 'O' is turned on in out-of-range for environmental operation condition.

Note: When the AED with displaying status indicator 'X' and temperature status is turned on in specified environmental operation condition, will operate properly.

Note: When the AED is turned on in the inappropriate environmental operation condition, temperature status will be blink.

Note: If the AED is placed in out-of-range value for environmental operation condition for long time, it will be longer than usual to recognize the temperature. It is recommended that AED should be stored in environmental operation condition described in this manual.

Note: If the AED was stored in low temperature conditions outside the operation environmental conditions for long time, place the AED in ambient temperature (20 °C) at least 0.5 hours for intended use. If the AED was stored in high temperature conditions outside the operation environmental conditions for long time, place the AED in ambient temperature (20 °C) at least 1 hours for intended use.

Install 1

- 1. To open the cover, push the Slide button to the right.
- 2. Check the pads are connected to the AED.
- If the pads are not connected to the AED, plug pads into the pads connector of the the AED.
- 4. Close the cover of the AED.
- 5. Install the battery to the AED.
- 6. The status indicator of the AED will display "X" and then operate the battery insertion self test.
- 7. When the battery insertion self test is completed normally, voice prompt "Unit ok" will be emitted and the status indicator will be changed from "X" to "O".

Note: The pads should be connected to the AED as preparation for emergency circumstances.

Note: Do not open defibrillation pads protective packaging until the time of emergency use when they are applied to a patient.

Note: If the battery insertion self-test is performed when the pads are not connected to the AED, the status indicator does not change from 'X' to 'O'.

Note: If 'X' is displayed on the status indicator even if the battery insertion self-test is proceeded with the pads connected to the AED, please contact qualified service personnel or your local supplier.

Install 2

Check that the AED is working optimally.

- 1. Change the Patient mode switch by pushing Slide button to the right or left for distinguish between adult and pediatric (infant-child).
- 2. Turn on the AED by pushing **Slide button** to the right and opening the cover, ensure that you can hear the voice prompt.
 - "Unit ok"
 - "Adult pads" or "Pediatric pads"
- 3. Ensure you can see the status indicator displays "O".
- 4. Turn off the AED by closing the cover.
- 5. Close the Cover with placing the defibrillation pads inside the AED.

Note: When the battery is replaced with the AED, self test will be automatically started. After completing the self test, ensure that you can hear the voice prompt "Unit ok" and then check the AED is turned off.

Install 3

Place the AED into its Soft Carry Case.

Install 4

Put into a storage or safe visible location.

Note: Storage differs in some countries. Ask qualified service personnel or your local supplier.

The AED should be kept in a convenient central area. Place it near a telephone so that the rescuer can call Emergency Medical Services and retrieve the AED without wasting time. Some important points to remember when storing:

- Store the AED in a suitable location for easy access.
- The AED should be placed in an appropriately accessible location.
- Store the AED in a clean and dry environment.
- Install the AED in the environmental operation condition described in this manual.

Make all necessary arrangements to ensure that the AED is accessible at all times. Inform any possible users of the location of the AED.

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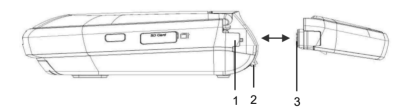
BATTERY OPERATION

⚠ WARNING	Test battery regularly, when the voltage of battery is very low. A battery that does not pass its test might shut down expectedly.
⚠ WARNING	Do not use a battery that is damaged, leaking, or wet.
⚠ WARNING	Do not use or store the battery in a place that may be exposed to high temperature.
⚠ CAUTION	When the voltage of the battery is very low, it is a possibility of not operating.
⚠ CAUTION	If the battery shows any signs of damage, leakage or cracking, it must be replaced immediately.
⚠ CAUTION	Discarded batteries may explode during incineration. Dispose used batteries properly. Do not dispose of batteries in refuse containers.
⚠ CAUTION	Check battery capacity regularly. Replace the new battery if you need.
⚠ CAUTION	Except for inspection, if the AED is frequently turned on, turned off or discharged, battery standby life will not last longer than the intended standby life by manufacturer.

Operating the AED on Battery Power

The AED has an installed non-rechargeable battery. The battery status appears on the status indicator when the AED is on battery power. The battery of AED is the handle part and replace the new handle/battery if need arises.

Replacing the battery



- 1. AED connector
- 2. Handle/Battery detachable button
- 3. Handle/Battery connector

Figure 4. Replacing the handle/battery - Right Panel

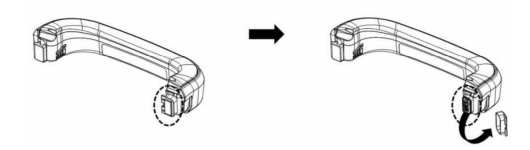
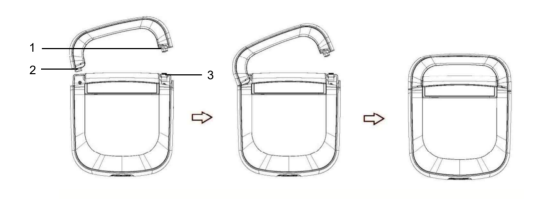


Figure 5. Removing the battery protection cap

Note: For new batteries, remove the battery protection cap before inserting the battery as shown in the Figure 5.



- 1. Handle/Battery connector
- 2. Hook
- 3. AED connector

Figure 6. Replacing the handle/battery - Upper Panel

- 1. Pull up while pushing the handle/battery detachable button, then disconnect the handle/battery.
- 2. Tilt up the handle/battery and then keep the handle/battery detachable button and AED connector perpendicular as shown in the Figure 4.
- 3. Connect the AED and handle/battery by using the hook as shown in the first figure of the Figure 6.
- 4. With the handle/battery being connected to the AED by hook, connect handle/battery connector and hook of the AED as shown in the second figure of the Figure 6.
- 5. When the both of the connection parts are fastened properly, the clink sound will be emitted.

The AED uses the non-rechargeable battery. Used battery is changed to new battery. Before turning on the AED with a battery that has been completely discharged, first replace the battery. When the new battery is installed, the AED is automatically turned on and then starts the battery insertion self test. After the battery insertion self test is completed, the AED may then be powered off.

Battery Status Indication

A new battery's life time is as below;

- Shelf life (in the original packaging): 2 years from manufacture date when stored and maintained according to direction provided in the operator's manual.
- Standby life (inserted in the AED): 5 years from manufacture date when stored and maintained according to direction provided in the operator's manual.
- Discharge: A minimum of 200 shocks (excepting the CPR period between the defibrillation therapies) or more than 6 hours of operating time under the ambient temperature at 20 °C.

If shock is delivered or shockable rhythm analysis is conducted once or more, the standby battery life would get shorter than the life time specified in above.

Note: After 200 times of shock, the voice prompt "Low battery, insert fresh battery" will be emitted.

Note: Due to the physical dimensions of the battery compartment, only batteries supplied by Mediana should be used. Using other types of replacement batteries may result in damage to the AED and void the limited warranty.

When operating on batteries, the battery status in the status indicator indicates the battery condition. See Table 4.

Table 4. The battery Status Icon

Battery Status Icons	Battery Status
	full charged (≤ 200 shocks or more than 6
	hours of operating time)
	used
	used (≤ 9 shocks)
	discharged (no shock)

If you hear the voice prompt "low battery, insert fresh battery" when the AED is turned on, the AED would be available 9 shocks. If the last bar of the battery indicator is invisible, buzzer would be sounded 2 times and then turned off automatically.

Self Test

Before using the AED, confirm that the AED is working properly and is safe to use as described below.

⚠ WARNING If the self test is not completed successfully, do not try to use the AED.

When power is applied, the AED automatically starts the self test, which tests the AED circuitry and functions. During performing Power On Self Test(POST), confirm that the AED status indicator turns on. If the AED status indicator does not function properly, do not use the AED. Instead, contact qualified service personnel or your local supplier.

Performing Power On Self Test (POST)

- 1. Turn on the AED by pushing the *Slide button*.
- 2. The AED automatically starts the Power On Self Test (POST).
- 3. If the AED detects an error during POST, the status indicator will display "X". Contact qualified service personnel or your local supplier for assistance.
- 4. Upon successful completion of the POST, the AED sounds voice prompt "Unit ok" and the status indicator displays "O".
- 5. Turn off the AED by closing the cover.

Automatic Self Test

The AED includes an automatic self test which is performed on a daily basis. The self test will run automatically and requires no user interaction. If there is an error, the status indicator displays "X".

The self test will test your AED and ascertain if its basic functions are running.

- Daily self test: MCU and Memory(RAM, ROM) integrity, Battery capacity, SD card connection and Patient body impedance.
- Weekly self test: Waveform delivery circuit low (2J) energy test, ECG circuit test in addition to the daily self test.
- Monthly self test: Waveform delivery circuit high (50J) energy test in addition to the weekly self test.

Note: When the battery is discharged, the status indicator will display "X". Even if the new battery is replaced, the status indicator still displays "X". Please contact qualified service personnel or your local supplier.

Note: Self test is not able to determine if the battery and the pads currently inserted in the AED are within their use by date. You must remember to check the use by date on the pads and standby life on the battery regularly.

Note: If automatic self-test is performed when the pads are not connected to the AED, the status indicator does not change from 'X' to 'O'.

Note: If 'X' is displayed on the status indicator even if automatic self-test is proceeded with the pads connected to the AED, please contact qualified service personnel or your local supplier.

Note: The internal battery is used to perform self-test and real time clock and it could operate for at least 10 years. However, please note that this period could vary

slightly depending on storage condition and use of environment. If the operating time has passed, the battery status disappears and "X" appears on the status indicator. In this case, please contact qualified service personnel or your local supplier.

Battery Insertion Self test

When the battery is installed or replaced, the AED automatically starts the battery insertion self test. After battery insertion self test is completed, the AED sounds voice prompt "Unit ok", the status indicator displays "O" and power of the AED is automatically turned off. If the battery insertion self test is not completed successfully, the AED sounds voice prompt "Unit fail" and the status indicator displays "X". If the AED does not function properly, do not use the AED. Instead, contact qualified service personnel or your local supplier.

You can also skip the battery insertion self test, try following procedure

- Closed: skip by opening the cover.
- Opened: skip by pressing the shock button.

After finishing this procedure, the AED will perform the power on self test as when user turns on the AED.

Note: Self test is not able to determine if the battery and the pads currently inserted in the AED are within their use by date. You must remember to check the use by date on the pads and standby life on the battery regularly.

Note: If the battery insertion self-test is performed when the pads are not connected to the AED, the status indicator does not change from 'X' to 'O'.

Note: If 'X' is displayed on the status indicator even if the battery insertion self-test is proceeded with the pads connected to the AED, please contact qualified service personnel or your local supplier.

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USING THE AED

⚠ WARNING	The AED should not be used on someone who is responsive when shaken or breathing normally.
⚠ WARNING	Do not use the pads if the pad gel is dried or damaged.
⚠ WARNING	Disconnect non-defibrillation protected electronic devices or equipment from patient before defibrillation.
⚠ WARNING	Never lift the AED by the pads cable or any other accessory. Such accessories could detach, causing the AED to fall on the patient.
⚠ CAUTION	Prolonged or aggressive CPR to a patient with pads attached can damage the pads. Replace the pads if they are damage during use or handling.

The AED is designed for the treatment of sudden cardiac arrest (SCA). It should only be used to treat someone who may be a victim of a SCA and is:

- · Unresponsive,
- · Non-breathing,
- Pulseless, (health care provider only)

If the person is unresponsive but you are unsure that they have suffered from a SCA begin CPR. When appropriate apply the AED and follow the voice prompts.

ERC guidelines (Rescue protocol)

The AED rescue protocol is consistent with the guidelines recommended by the ERC 2021 Guidelines for Resuscitation and Emergency Cardiac Care. The AED rescue protocol is subject to be upgradeable in order to be consistent with and optimized for the guidelines recommended by the latest version of ERC Guidelines for Resuscitation and Emergency Cardiac Care. Please contact your Mediana service representative for more information.

Note: ERC is the abbreviation for 'European Resuscitation Council'.

Note: This section is described in accordance with ERC Guidelines.

Summary of CPR Guidelines

This "Guidelines Highlights" publication summarizes the ERC 2021 Guidelines. This is easy reference material for both lay rescuer and healthcare provider.

Ensure scene safety.	Make sure you, the victim and any bystanders are safe.		
Check the victim for a response.	Gently shake victim's shoulders and ask loudly: "Are you all right?"		
Open the airway, Check for	Open the airway.		
breathing.	Look, listen and feel for normal breathing.		
	Person not responsive?		
	Not breathing normally?		
	Ask a helper to call the emergency services if possible otherwise call them yourself.		
After call for help, Send for AED.	Stay with the victim when making	the call if possible.	
	Send someone to find and bring an AED if available.		
	If you are on your own, do not leave the victim, start CPR.		
	•		
	Start chest compressions.		
30 Compressions 2 Breaths UNTIL EMC ARRIVE.	After 30 compressions open the airway again and deliver two rescue breaths.		
	Note: If unable to do rescue breaths, give chest compressions only CPR.		
	Continue CPR until an AED is available or arrival of emergency physician.		
	If the AED available, turn on and	follow instructions.	
	ANALYSIS SHOCK DECISION		
	YES	NO	
Repeat every 2 minutes.	1		
	DELIVER SHOCK.		
	1	•	
	Continue CPR for about 5 cycles. (approximately 2minutes)		

Pre Defibrillation Action

Prior to using the AED, it is advised to perform the following checks and actions in order to prepare the patient.

- Remove clothes to expose bare chest.
- If excessively hairy shave hair from areas to which defibrillation pads are to be applied.
- Ensure that the patient chest is dry. If necessary, dry chest area.

Operating the AED

↑ WARNING	The Pediatric (Infant-Child) mode must be used on patients between 1
<u> </u>	and 8 years or less than 25 kg (55lb).
⚠ WARNING	If the pads placement is inappropriate, the AED could harm the patient. To place the accurate position, must follow the voice prompt and action icon. When pads placement is inappropriate, treatment could not work or shock could burn the patient's skin.
⚠ WARNING	Do not place pads near the generator of an internal pacemaker. The analyzing heart rhythm of patient who is implanted pacemaker could inaccurate or the pacemaker might be damaged by defibrillator discharges.
⚠ WARNING	Do not perform chest compressions (CPR) through electrodes. These actions may damage the electrode pads cause the AED to function improperly. If CPR is performed on a child with pads attached in anterior posterior, check the pads for damage and make sure the pads are properly attached after performing CPR.
⚠ WARNING	Always apply pads to flat areas of skin. Avoid application over folds of skin such as those underneath the breast or on obese patients. Excessive hair, poor adhesion, or air under pads may produce burns or ineffective energy transfer.
⚠ WARNING	To apply the pads to patient chest properly, shave hair from areas which defibrillation pads are to be applied if necessary.
⚠ WARNING	Always check the use by date on the pads and do not use the pads if the packaging has been previously opened. If the excessively dry pads are attached, AED may interpret as a condition that the pads are not attached to the patient.
⚠ WARNING	Touching the patient during the analysis phase of treatment can cause interference with the diagnostic process. Avoid contact with the patient while analysis is being carried out. The AED will instruct you by voice prompt when it is safe to touch the patient.
⚠ WARNING	The AED delivers shocks which can cause serious harm to operators and bystanders. Caution must be taken to ensure no-one is in contact with the patient when a shock is delivered.

Note: Only pads supplied by Mediana should be used. Using other types of pads may result in damage to the patient and the AED.

Note: If the AED is turned on in out-of-range of environmental operation condition described in this manual, temperature status will be blinked. In this case, place the AED in appropriate temperature before use it.

- 1. Check the status indicator displays "O".
- 2. To open the cover, push the Slide button to the right.
- 3. Turn on the AED by opening the cover.
- 4. The AED automatically starts the Power-On-Self Test.
- 5. The test result is displayed on the status indicator and the voice prompt sounds.
 - Self test is passed : Voice prompt "Unit ok", Status indicator "O"
 - Self test is failed: Voice prompt "Unit failed", Status indicator "X"
- 6. If the pads are inserted and the Patient mode switch is selected, you will hear the voice prompt:
 - Patient mode switch is switched to left, "Adult pads"
 - Patient mode switch is switched to right, "Pediatric pads"

If the pad is not inserted, you will hear the voice prompt:

• "Plug in pads. Insert connector firmly."

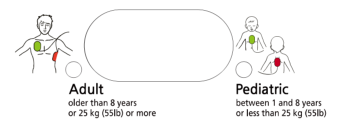


Figure 7. Patient mode switch

Note: The patient mode can switch even if any step except for CPR is going on progress. If the patient mode is changed, the AED will emit the voice prompt "Adult pads" or "Pediatric pads". When the patient mode is changed during CPR, the AED will not emit the voice prompt.

Note: Return to ECG analysis, while analyzing ECG or delivering electric shock, even though the patient mode is changed during process. When the number of mode change is 3 times, the AED will proceed the CPR.

7. Verify the AED up to '6.' which is activated normally and follow voice prompt and action icon. The red LED will flash under the relevant action icon.

Operation of HeartOn A15

Note: Follow voice prompt. Do not touch patient or allow any others to touch the patient while the AED is analyzing. After completion of analysis, the AED will advise you of treatment recommended. Care must be taken to keep the patient still. A moving patient can lead to incorrect, delayed or less effective diagnosis and therapy.

Note: The AED will only administer a shock if it is needed. A voice prompt will tell you when to press the shock button to administer defibrillation therapy.

Note: If you hear below voice prompt when the AED is turned on, the following actions should be performed:

• "Low battery, insert fresh battery."

After inserting the new battery, open the AED cover to turn on the AED. The power on self-test will be performed automatically.

Step 1

• "Check for response. Are you all right?"



Figure 8. HeartOn A15: Action Icon - Step 1

Step 2

• "Call for help."



Figure 9. HeartOn A15: Action Icon - Step 2

Step 3

• "Open the airway."



Figure 10. HeartOn A15: Action Icon - Step 3

Note: ERC 2021 Guidelines indicates that user should open the airway before checking for breathing.

Step 4

• "Check breathing."



Figure 11. HeartOn A15: Action Icon - Step 4

Step 5

Remove clothes to expose the patient's chest. If the patient has an excessively hairy chest, shave the area where the pads are about to be applied.

• "Remove clothes from the patient's chest. Place pad exactly as shown in the picture. Press pads firmly to patient's bare chest."

When pads are disconnected to the AED, the following voice prompt will be emitted.

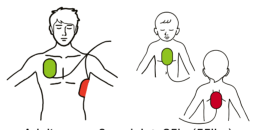
• "Plug in pads. Insert connector firmly."

When the pads are connected to the AED according to the voice prompt, the following voice prompt will be emitted.

• "Adult pads" or "Pediatric pads"



Figure 12. HeartOn A15: Action Icon - Step 5



Adult: age > 8, weigh \geq 25kg(55lbs) Pediatric (Infant-Child): age \leq 8, weigh \leq 25kg(55lbs)

Figure 13. Pads Placement

⚠ WARNING

Apply freshly opened and undamaged pads, within use by date, to clean and dry skin to minimize burning.

Step 6

When the pads are attached correctly to the patient you will hear the voice prompts:

- "Do not touch the patient."
- "Analyzing and Charging"
- "Shock advised."
- "Do not touch the patient."
- "Analyzing and Charging"
- "No shock advised."

When the patients' ECG rhythm changes to a non-shockable rhythm, when the pads are disconnected, when the impedance changes, or the patient mode is changed, analyzing and charging are suspended and the following voice prompt will be emitted:

"Shock cancelled."



Figure 14. HeartOn A15: Action Icon - Step 6

Note: If "No shock advised", the AED will move to step 8 which demonstrate CPR process directly.

Note: The AED performs the Step 6 directly when it is turned on after the rescuer attaches the pads to the patient properly. Also, the Step 6 would be started if the pads are attached to the patient even if the AED is under the Step 1 to 5. This can reduce the preparing time for electric shock.

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Step 7

- "Press the flashing button now." {Repeat twice}
- "Shock delivered."

When the shock button is not pressed, the following voice prompt will be emitted:

- "Press the flashing button now." {Repeat twice}
- "Shock button not pressed."

When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed, the following voice prompt will be emitted:

"Shock cancelled."



Figure 15. HeartOn A15: Action Icon - Step 7

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Note: If "Shock button not pressed.", the AED will move to step 8 which demonstrate CPR process directly.

Step 8

- "It is safe to touch the patient."
- "Begin CPR." {Beep, (30 times at a rate of 120/min)} or "If needed, Begin CPR." {Beep, (30 times at a rate of 120/min)}
 - "Give two breaths." {Repeat 5 times}
- "Stop CPR."
- "Adult pads" or "Pediatric pads"



Figure 16. HeartOn A15: Action Icon - Step 8

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Note: After finishing Step 8, the AED will move to the Step 6 to analyze the heart rhythm again. The voice prompt for the type of pads will be emitted.

Operation of HeartOn A15-G4

Note: Follow voice prompt. Do not touch patient or allow any others to touch the patient while the AED is analyzing. After completion of analysis, the AED will advise you of treatment recommended. Care must be taken to keep the patient still. A moving patient can lead to incorrect, delayed or less effective diagnosis and therapy.

Note: The AED will only administer a shock if it is needed. A voice prompt will tell you when to press the shock button to administer defibrillation therapy.

Note: If you hear below voice prompt when the AED is turned on, the following actions should be performed:

"Low battery, insert fresh battery."
 After inserting the new battery, open the AED cover to turn on the AED. The power on self-test will be performed automatically.

Connection of the pads

When the pads are not connected during the AED operation, the following voice prompt will be emitted.

"Plug in pads. Insert connector firmly."

When the pads are connected to the AED according to the voice prompt, the following voice prompt will be emitted.

• "Adult pads" or "Pediatric pads"



Figure 17. HeartOn A15-G4: Pad disconnect icon

Note: If the pad connector is not connected in any step except for Step 4 (CPR), the AED will move to Pad connector disconnected icon and the voice prompt "Plug in pads. Insert connector firmly." is emitted.

Step 1

Remove clothes to expose the patient's chest. If the patient has an excessively hairy chest, shave the area where the pads are about to be applied.

• "Remove clothes from the patient's chest. Place pad exactly as shown in the picture. Press pads firmly to patient's bare chest."



Figure 18. HeartOn A15-G4: Action icon - Step 1



Adult: age > 8, weigh ≥ 25 kg(55lbs) Pediatric (Infant-Child): age ≤ 8 , weigh < 25kg(55lbs)

Figure 19. Pads placement

Step 2

When the pads, as shown in the figure 19 are attached correctly to the patient you will hear the voice prompts:

- "Do not touch the patient."
- "Analyzing and Charging"
- "Shock advised."
- "Do not touch the patient."
- · "Analyzing and Charging"
- "No shock advised."

When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed, analyzing and charging are suspended and the following voice prompt will be emitted:

"Shock cancelled."



Figure 20. HeartOn A15-G4: Action icon -Step 2

Note: If "No shock advised" voice prompt is emitted, the AED will move to step 4 which demonstrate the CPR process.

Note: The AED performs the Step 2 directly when it is turned on after the rescuer attaches the pads to the patient properly. Also, the Step 2 would be started if the pads are attached to the patient even if the AED is under the Step 1. This can reduce the preparing time for electric shock.

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Step 3

- "Press the flashing button now." {Repeat twice}
- "Shock delivered."

When the shock button is not pressed, the following voice prompt will be emitted:

- "Press the flashing button now." {Repeat twice}
- "Shock button not pressed."

When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed, the following voice prompt will be emitted:

"Shock cancelled."



Figure 21. HeartOn A15-G4: Action Icon - Step 3

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Note: If the voice prompt "Shock button not pressed" is emitted, the AED will move to Step 4 to perform CPR.

Step 4

- "It is safe to touch the patient."
- "Begin CPR." {Beep, (30 times at a rate of 120/min)} or "If needed, Begin CPR." {Beep, (30 times at a rate of 120/min)}
 - "Give two breaths." {Repeat 5 times}
- "Stop CPR."
- "Adult pads" or "Pediatric pads"



Figure 22. HeartOn A15-G4: Action Icon - Step 4

Note: If the shock cancel (When the patients' ECG rhythm is changed to a non-shockable rhythm, when the pads are disconnected or when the impedance or mode is changed) has been occurred 3 times, the AED is performing the CPR process.

Note: If the pads connector has been disconnected in Step 4 (CPR), the AED moves to the pad disconnect icon after the CPR. If the pads are still connected, the AED moves to Step 2 to analyze the heart rhythm again and the type of currently attached pads is announced through a voice prompt.

Performing CPR

After the electric shock is delivered, the following voice prompt would be emitted.

- "It is safe to touch the patient."
- "Begin CPR."

Follow the voice prompts to properly perform the CPR.

When performing CPR, use the metronome sound from the AED for compression rate – the AED emits a tone at a rate of 100 beats per minute. Also, action icon of the AED will be flashing at a same rate of metronome sound.

Rescuer performs 5 cycles of CPR, each cycle includes 30 times of chest compression and 2 times of rescue breaths. Or perform the chest compression without rescue breath, if unable to do rescue breaths. The AED will remain in CPR mode for 5 cycles (approximately 2 minutes). After CPR mode you will hear the following voice prompt:

• "Stop CPR."

The AED will then return to ECG analyzing procedure. Continue to follow this instruction until emergency physician arrives and then hand over patient to emergency physician.

Note: In accordance with ERC 2021 Guidelines,

- the recommended compression rate is 100 ~ 120 beats per minute,
- the recommended compression depth is at least 2 inches (5 cm), but not more than 2.4 inches (6 cm),
- the recommended compression ventilation ratio is 30:2,
- the recommended duration is 5 cycles (30:2 x 5 cycles).

Note: In all cases follow the voice prompts and visual instructions given by the AED and/or the emergency services. You can find more information about CPR at the following links:

- https://www.nhs.uk/conditions/first-aid/cpr/
- https://www.resus.org.uk/home/faqs/faqs-basic-life-support-cpr

Note User and Bystander Safety



Make sure no one is touching the patient before you press the Shock button. Loudly announce, "Stand back! Do not touch the patient." And look down the entire length of the patent to ensure there is no contact before pressing the Shock button.

Do not touch the patient while the AED is analyzing or delivering a shock is in process. Defibrillation energy can cause injury. As long as the AED is used according to the directions, and no one is in contact with the patient when the **Shock button** is pressed, there is no risk of harm to the rescuer or bystanders. The AED cannot deliver a shock unless the pads are applied to someone whose heart is in need of a shock.

Note: See warnings and cautions for more details.

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MAINTENANCE

	Improper maintenance which is provided in this manual may damage the
⚠ WARNING	AED or cause it to function improperly. Maintain the AED according to
	directions.
↑ WARNING	Do not let fluids to get into the AED. Avoid spilling any fluids on the AED
	or its accessories. Spilling fluids into the AED may damage it or cause a
	fire or electric shock hazard. Do not sterilize the AED or its accessories.
	Do not immerse any part of the AED in water or any type of fluid. Contact
⚠ WARNING	with fluids may seriously damage the AED or cause fire or electric shock
	hazard.
↑ WARNING	Do not attempt to warm the electrodes with a heat source greater than
	35℃ (95°F).
⚠ WARNING	Do not clean the AED with abrasive materials, cleaners or solvents.
↑ CAUTION	Follow local government ordinances and recycling instructions regarding
CAUTION	disposal or recycling of AED components, including batteries.
	Do not short-circuit the battery, as it may generate heat. To avoid short-
⚠ CAUTION	circuiting, do not let the battery terminal come in contact with metal
	objects at any time, especially when transporting.
⚠ CAUTION	Do not solder the battery directly. Heat applied during soldering may
	damage the safety vent in the battery's positive cover.
⚠ CAUTION	Do not deform the battery by applying pressure. Do not throw, hit, drop, fold or impact the battery.
	Do not use the battery with other maker's batteries, different types or
⚠ CAUTION	models of batteries such as dry batteries, nickel-metal hydride batteries,
CAUTION	or Li-ion batteries together, as they might leak electrolyte heat or explode.
^	Do not mistreat the battery, or use the battery in applications not
⚠ CAUTION	recommended by Mediana.
A CALITICAL	Keep the battery out of reach of babies and children to avoid any
⚠ CAUTION	accidents.
↑ CAUTION	If there are any problems with the battery, immediately put the battery in a
ZI CAUTION	safe place and contact qualified service personnel or your local supplier.
⚠ CAUTION	Replacing new battery and placing the pads should carry out in
	environmental conditions described in this manual. If the AED is operated
	in out-of-range for environmental conditions, the AED can't be operated
	properly.

After using the AED, Mediana technical support recommend you perform the following actions:

- 1. Use the HeartOn AED Event Review Software to download information about the therapy performed and store appropriately. (If you do not have the HeartOn AED Event Review Software, please contact your dealer who can arrange for the incident to be downloaded)
- 2. Remove the used the pads from your AED and dispose of in a suitable manner. (For recommended disposal methods please refer to section the recycling and disposal)
- 3. Check the exterior of the AED for cracks or other signs of damage. Contact your distributor or Mediana technical support immediately if any damage is found.
- 4. Check the exterior of the AED for dirt or contamination. If necessary, clean the AED with approved cleaning products.
- 5. Check supplies, accessories and spares for damage or expiration. Replace immediately if any damage or expiration is found. Contact your local Mediana approved dealer.

- Install the new pads or battery. Before installing the new pads check that its use by date has not been exceeded.
- 7. After installation of the new battery. Check the status Indicator. If the status Indicator is not displaying "O" refer to the troubleshooting section of this manual. If the problem persists, contact Mediana or your local approved dealer for technical support.
- 8. Turn on the AED and verify that the AED operates in the correct manner i.e. voice prompt "Unit OK" can be heard. Turn off the AED.
- 9. Contacting Mediana after use. At Mediana we like to hear from our customers whenever they have any occasion to use any of our products, even if therapy is not delivered as part of the incident. This information is vital to the continued development and constant improvement we strive for in the treatment of sudden.

Recycling and Disposal

When the AED, battery or accessories reach the end of useful life, recycle or dispose of the equipment according to appropriate local and regional regulations.

Note: The AED should be disposed of separately from the municipal waste stream via designated collection facilities appointed by the government or the local authorities.

Note: The correct disposal of your old appliance will help prevent potential negative consequences for the environment and human health.

Note: For more detailed information about disposal of your old appliance, please contact your city office, waste disposal service or the shop where you purchased the AED.

Returning the AED and System Components

To return the AED and/or accessories, contact qualified service personnel or your local supplier.

Service

The AED requires no routine service other than cleaning, battery maintenance, and service activity which is mandated by the user's institution. For more information, refer to the AED service manual. Qualified service personnel in the user's institution should perform periodic inspections of the AED. If service is necessary, contact qualified service personnel or your local supplier.

Periodic Safety Checks

It is recommended that the following checks be performed every month.

- Inspect the equipment for mechanical and functional damage.
- · Inspect the external safety labels for legibility.

Cleaning

To clean the AED, wipe the AED with a soft cloth that has been dampened by one of the following:

- · Soapy water.
- Isopropyl alcohol (70% solution).

For cables and pads, follow cleaning instructions in the directions for use shipped with those components.

Avoid spilling liquid on the AED, especially in connector areas. If liquid is accidentally spilled on the AED, clean and dry thoroughly before reuse. If in doubt about AED safety, refer the unit to qualified service personnel or your local supplier for checking.

Battery Maintenance

The new battery lifetime in use can be more than 6 hours monitoring or 200 shocks (excepting the CPR period between the defibrillation therapy) or a combination of both. The battery in the standby mode (inserted into the AED) has standby life (5 years from manufacture date). If the battery status is flashing with one bar, you may need to replace the battery with a fresh battery. If the battery is not inserted into the AED, the battery has a shelf life. (2 years from manufacture date)

For diagnosis of the reason for status indicator display "X", please refer to the troubleshooting section.

Pads Maintenance

Replacement of the pads must be carried out if:

- The use by date of the pads has been exceeded.
- When the pads have been used. (It is a single use item and must be replaced with new pads.)
- The package of new pads has been previously damaged.
- The pads have been vent.

Replacing Pads

- 1. Take the replacement pads from its protective bag.
- 2. Disconnect the pad connect from the AED.
- 3. Push the pads firmly to ensure it is fully inserted.
- 4. Turn on the AED.
- 5. Check the status indicator. If the pads have been inserted correctly, the status Indicator displays "O".
- If necessary inform relevant safety officer or person responsible for maintenance of the AED.
- 7. Update the relevant information to show the date that the replacement of pads and battery was placed into service.
- 8. Dispose of the old pads.

The AED Maintenance

Mediana recommends users perform regular maintenance checks. A suggested maintenance check would be.

- 1. Check the status Indicator. If the status Indicator displays "X", a problem has been detected. Refer to the troubleshooting section of this manual.
- 2. Check the use by date of the pads. If the pads have exceeded its use by date , remove it and replace with the pads. Contact qualified service personnel or your local supplier for replacements.



Figure 23. Use by date of Pads

- 3. Check the AED and accessories for damage or use by date. Replace any accessories found to be damaged or that have exceeded their use by date.
- 4. Check the exterior of the AED for cracks or other signs of damage. Contact qualified service personnel or your local supplier if any damage is found.
- 5. Check that building users are aware of AED location and that it is easily accessible at all times.

TROUBLESHOOTING

⚠ WARNING	If you are uncertain about the accuracy of any measurement, check the patient's vital signs by alternate means; then make sure the AED is functioning correctly.
⚠ WARNING	To reduce the risk of electrical shock, do not attempt to remove the cover under any circumstances. There are no operator serviceable components and only a qualified technician should service the AED.

General

If the AED detects an error, it can display the "X" on the status indicator. Contact qualified service personnel or your local supplier. Before calling to qualified service personnel or your local supplier, make sure it meets environmental conditions provided in the manual as temperature, humidity, altitude and so on.

Corrective Action

Check used by date the pads. Change the pads if use by date has been exceeded. Check shelf life or standby life of the battery. Change the battery if the shelf life or standby life has been exceeded.

Following is a list of possible errors and suggestions for corrective action.

If the status indicator is still not displaying "X" or a warning message is heard when the AED is turned on or if for any reason, you have suspicions that your AED is not working correctly contact qualified service personnel or your local supplier or Mediana directly for support. (info@mediana.co.kr)

1. There is no response to the opening the cover of the AED.

- A CPU module may be malfunctioned. Notify qualified service personnel or your local supplier to check and replace the CPU module.
- The battery may be missing or discharged. If the battery is missing, insert the battery (See Battery Operation section). If the battery is discharged, change the battery. (See Maintenance section)

2. The beep tones do not sound during the operation.

- Do not use the AED; contact qualified service personnel or your local supplier.
- 3. The beep tones sound but voice does not function properly.
 - Do not use the AED; contact qualified service personnel or your local supplier.
- 4. The voice prompt "Plug in pads. Insert connector firmly".
 - Reconnect the pad connector with pad socket firmly or replace the pad.
- 5. The action icon does not flash.
 - Do not use the AED, contact qualified service personnel or your local supplier.
- 6. The voice prompt is unclearly heard.
 - Do not use the AED, contact qualified service personnel or your local supplier.
- 7. The battery status does not indicate 3 bar despite of replacing new battery.
 - If the battery status still not displayed 3 bar despite of replacing new battery, do not use the AED and contact qualified service personnel or your local supplier.

EMI (Electromagnetic Interference)

⚠ WARNING	Keep patients under close surveillance during delivering a shock. It is possible, although unlikely, that radiated electromagnetic signals from sources external to the patient and the AED can cause inaccurate measurement readings. Do not rely entirely on the AED readings for patient assessment.
⚠ WARNING	It is possible that any radio frequency transmitting equipment and other nearby sources of electrical noise may result in disruption in the AED operation.
⚠ WARNING	It is possible, although unlikely, that large equipment using a switching relay for its power on/off may affect the AED operation. Do not operate the AED in conjunction with electrocautery or diathermy equipment or in such environments.

This AED has been tested and found to comply with the limits for Medical devices to the IEC60601-1-2, and the Medical device Directive 93/42/EEC. These limits are designed to provide reasonable protection against harmful interference in a typical medical installation.

However, because of the proliferation of radio-frequency transmitting equipment and other sources of electrical noise in health care environments (such as electrosurgical equipment, cellular phones, mobile two-way radios, electrical appliances, and high-definition television), it is possible that high levels of such interference due to close proximity or strength of a source may affect AED operation.



The AED is designed for use in environments in which the signal can be obscured by electromagnetic interference. During such interference, measurements may seem inappropriate or the AED may not seem to operate correctly.

The AED disruption may be indicated by erratic readings, cessation of operation, or other incorrect functioning. If this occurs, survey the site to determine the source of this disruption. Try the following actions to see if they eliminate the disruption:

- Turn equipment in the vicinity off and on to isolate the offending equipment.
- Reorient or relocate the interfering equipment.
- Increase the separation between the interfering equipment and this equipment.

The AED generates, uses, and can radiate radio frequency energy. If the AED is not installed and used in accordance with these instructions, the AED may cause harmful interference with other devices in the vicinity.

If assistance is required, contact qualified service personnel or your local supplier.

Obtaining Technical Assistance

For technical information and assistance, or to order the AED service manual, call your local supplier. The service manual provides information required by qualified service personnel or your local supplier when servicing the AED.

GLOSSARY

Sudden Cardiac Arrest (SCA)

Sudden cardiac arrest is a condition in which the heart suddenly stops pumping effectively due to a malfunction of the heart's electrical system. Often victims of SCA have no prior warning signs or symptoms. SCA can also occur in people with previously diagnosed heart conditions. Survival for an SCA victim depends on immediate cardio-pulmonary resuscitation (CPR). The use of an external defibrillator within the first few minutes of collapse can greatly improve the patients' chances of survival. Heart attack and SCA are not the same, though sometimes a heart attack can lead to a SCA. If you are experiencing symptoms of a heart attack (pain, pressure, shortness of breath, squeezing feeling in chest or elsewhere in the body) seek emergency medical attention immediately.

Heart Rhythm

The normal electrical rhythm by which the heart muscle contracts to create blood flow around the body is known as Sinus Rhythm. Ventricular Fibrillation (VF) caused by chaotic electrical signals in the heart is often the cause of SCA, but a shock can be administered to re-establish sinus rhythm. This treatment is called defibrillation. The AED is designed to automatically detect ventricular fibrillation (VF) and perform defibrillation on victims of sudden cardiac arrest.

Ventricular Tachycardia / Ventricular Fibrillation

Is a life-threatening heart rhythm that is treatable with the therapy using the AED.

Sinus Rhythm

Sinus Rhythm is the normal electrical rhythm by which the heart muscle contracts and expands to create blood flow around the body.

Biphasic Shock

A biphasic shock is an electrical current that is passed through the heart, firstly in one direction and then in another.

Biphasic Truncated Exponential (BTE) waveform

Biphasic Truncated Exponential (BTE) waveform stands for Self-Compensating Output Pulse Envelope Waveform.

Pads

Pads are the electrodes that are connected to the patient's chest in order to administer therapy.

Electromagnetic Interference

Electromagnetic interference is radio interference that may cause erroneous operation of electronic equipment.

Impedance Measurement

Impedance measurement is a check that is performed to check the integrity of AED patient contact.

Detecting Fibrillation

The electrical rhythm by which the heart muscle contracts can be detected and used for medical diagnosis and the resulting reading is called an Electrocardiogram (ECG). The AED has been designed to analyze a patient's ECG in order to detect ventricular fibrillation (VF) in the heart. If ventricular fibrillation (VF) is detected the AED will deliver a carefully engineered electrical shock designed to stop the chaotic electrical activity experienced within the heart muscle during SCA. This may allow the victim's heart to return to a sinus rhythm.

HeartOn A15

The AED is a semi-automatic device used for the delivery of external defibrillation therapy to resuscitate victims of SCA, who are unresponsive, are not breathing, or without life signs.

HeartOn AED Event Review Software

HeartOn AED Event Review Software is software that can be used in conjunction with the AED and SD card (or Infrared communication cable). It can retrieve and view information about therapy delivered using the AED. Also, HeartOn AED Event Review Software can be used to configure the AED.

More Information

If you have had any occasion to use your AED or if you require any further information on the AED, its accessories or any other products please contact us.

SPECIFICATION

Defibrillation Electric Shock

\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Biphasic Truncated Exponential (BTE) waveform
Waveform	(impedance compensation)
	HeartOn A15 / HeartOn A15-G4
Energy	Adult: 170 to 195J (±5%)
	Pediatric (Infant-Child): 44 to 51J (±5%)
Operating mode	Semi-Auto

ECG

Lead	II (RA, LL)		
Patient impedance	HeartOn A15 / HeartOn A15-G4 25 to 200 ohm		
	Note: Patient impedance depends on the customer		
	request.		
Heart Rate	20 to 300 per min		
Accuracy	1 per min		
Detection	V/F greater than or equal to 0.2 mV		
	V/T Adult: greater than or equal to 150 bpm		
	Pediatric (Infant-Child): greater than or equal to		
	180bpm		
Lead connection	Confirm the connection and emit voice prompt		
Filter	0.5 to 30 Hz		

Indication

Controls			
Standard	Slide button, Shock button, Patient mode switch		
	Indicators		
Visible	ICON Indicator, Status LCD(AED status, Battery status,		
	Temperature status), LED (Patient mode switch LED)		
Audible	Audio speaker (Voice prompt)		
	Beep (CPR indication, Power on, Critically Low or Low		
	Battery, Self-Test fail, Cover operation defects, Charging		
	time is too long)		

Physical

Dimensions	240 × 294 × 95 (mm) (W×H×D)
Weight	Approx. 2.5 kg including the battery excluding pads

Environmental Conditions

Operation/Standby		
Temperature	0 to 43°C (32 to 109.4°F)	
Relative Humidity	5 to 95% RH (N	on-condensing)
Altitude	0 to 4,575 m	
Shock	Acceleration:	100 G (+/- 10%)
	Time:	6 msec
	The number of	shocks: 3 times/axis (6 axes (+/- X, Y, Z))
Vibration	Frequency:	10Hz to 2000Hz
	Acceleration:	10 Hz to 100 Hz: 5,0 (m/s²)²/Hz
		100 Hz to 200 Hz: -7 dB per octave
		200 Hz to 2000 Hz: 1,0 (m/s²)²/Hz
Drop height	1m	
Water and dust	IP54 (IEC60529	9)
resistance		
Note: The Standby condition indicates that the self test periodically runs with		
installed battery in the AED.		
Storage (in shipping container)		
Temperature	-20 to 60°C (-4 to 140°F)	
Relative Humidity	5 to 95% RH (Non-condensing)	
Altitude	0 to 12,192 m	

Self Test

Self Test		
Temperature	0 to 43°C (32 to 109.4°F)	
Cycle	Every 24 hours, 1 week, 1 month	
	Power on self test, Battery insertion self test	
Test result Status LCD displays "O"/ "X".		
Note: Only when the battery is installed, self test will be activated.		

Data Backup and Communication

Standard SD card, Infrared communication port

Expected Service Life

Expected service life	10 years

Accessories Specifications

Pads

Adult / Pediatric (Infant-Child) Pads			
Shelf life	Refer to pad's direction for use		
Electrodes	Disposable pa	ds	
Placement	Adult: Anterior-lateral		
	Pediatric (Infant-Child): Anterior-posterior		
Minimum active gel	80 cm ² +/-5%		
area			
Cable length	About 1.8 m		
Environmental Conditions			
Temperature	Operation:	0 to 43°C (32 to 109.4°F)	
	Storage:	0 to 43°C (32 to 109.4°F)	
Relative Humidity	5 to 95% RH (Non-condensing)		

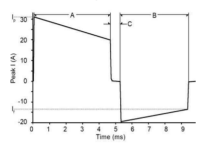
Battery

Battery				
Туре	LiMnO ₂ , Disposable, Long-Life Primary Cell			
Voltage/Capacity	15V, 4200 mAh	1		
Shelf Life (in the	2 years from m	anufacture date		
original packaging)				
Standby Life	5 years from m	anufacture date		
(inserted in the AED)				
Discharge	A minimum of 200 shocks (excepting the CPR period			
	between the defibrillation therapy) or more than 6 hours of			
	operating time at 20°C			
Environmental Conditions				
Temperature	Operation:	0 to 43°C (32 to 109.4°F)		
	Storage:	0 to 43°C (32 to 109.4°F)		
Relative Humidity	5 to 95% RH (Non-condensing)			

Defibrillation waveform







Waveform parameters are automatically adjusted as a function of patient defibrillation impedance. In the diagram at left, A is the width of pulse 1 and B is the width of pulse 2 of the waveform, C is the inter-pulse delay, I_p is the peak current, and I_f the final current.

The AED delivers shocks to load impedances from 25 to 200 ohm. The duration of each pulse of the waveform is dynamically adjusted based on delivered charge, in order to compensate for patient impedance variations, as shown below:

Adult defibrillation					
Load	Pulse width 1	Pulse width 2	Delivered		
Resistance (Ω)	(ms)	(ms)	Energy (J)		
25	3.3	3.1	195		
50	4.7	4.1	190		
75	6.7	4.7	185		
100	8.3	5.9	195		
125	9.7	6.7	190		
150	11.3	7.9	185		
175	11.7	8.7	180		
200	11.7	8.7	170		
Pe	Pediatric (Infant-Child) defibrillation				

rediatio (illiant-child) delibrillation				
Load	Pulse width 1	Pulse width 2	Delivered	
Resistance (Ω)	(ms)	(ms)	Energy (J)	
25	3.3	3.1	51	
50	4.7	4.1	50	
75	6.7	4.7	49	
100	8.3	5.9	51	
125	9.7	6.7	50	
150	11.3	7.9	49	
175	11.7	8.7	47	
200	11.7	8.7	44	

Charge control

Controlled by patient analysis system for automated operation.

Charging Time

< 10 seconds typical (with new battery)

Shock Analysis Time < 8 seconds typical

Note: Shock analysis would take about 16 seconds at maximum from the start to the completion of analysis if ECG signal is interfered by motion or others.

Disarm

After the AED advises a shock, AED continues to monitor the patient ECG rhythm during a maximum of 20 seconds until the shock button is pressed. If the ECG rhythm changed to a nonshockable rhythm, the shock energy that is accumulated on the high-voltage capacitor will be disarmed through internal resistance. If the shock button is not pressed during 20 seconds, the shock energy that is accumulated on the high-voltage capacitor will be disarmed through internal resistance.

ECG Analysis Performance

Rhythm class	AHA-DB	MIT-DB	CU-DB	VF-DB	VT-DB	Total number of sample size
VF-shockable : TP	988	22	282	908	-	2200
VF-shockable : FN	44	0	7	23	-	74
VF-shockable : sensitivity(%)	95.74	100.00	97.58	97.53	-	96.75
Adult mode VT-shockable : TP	-	-	-	701	-	701
Adult mode VT-shockable : FN	-	-	-	66	-	66
Adult mode VT-shockable : sensitivity(%)	-	-	-	91.40	-	91.40
Pediatric (Infant-Child) mode VT-shockable : TP	-	-	-	-	18	18
Pediatric (Infant-Child) mode VT-shockable : FN	-	-	-	-	3	3
Pediatric (Infant-Child) mode VT-shockable : sensitivity(%)	-	-	-	-	85.71	85.71
Non-shockable : TN	32291	17518	1267	7062	-	58138
Non-shockable : FP	0	2	8	33	-	43
Specificity(%)	100.00	99.99	99.37	99.53	-	99.93
Positive Predictive Value	100.00	91.67	97.24	97.99	100.00	98.55

Database for ECG Analysis

- From AHA (American Heart Association) official database
- From MIT (Massachusetts institute Technology) official database (CU-DB: The Creighton University Sustained Ventricular Arrhythmia Database, VF-DB, VT-DB: MIT-BIH Malignant Ventricular Arrhythmia Database)

ECG rhythm to determine if a shock is appropriate

- Ventricular Fibrillation at a amplitude greater than or equal to 0.2mV.
- Ventricular Tachycardia at a heart rate greater than or equal to 150 bpm (Adult) / 180 bpm (Pediatric (Infant-Child)).

Compliance

ltem	Standard	Description
Classification	IEC60601-1:2005	Internally powered (on battery power)
	+A1:2012+A2:2020	
	EN 60601-1:2006	
	A1:2013+A2:2021	
Type of	IEC60601-1:2005	Type CF – Applied part
protection	+A1:2012+A2:2020	
	EN 60601-1:2006	
	A1:2013+A2:2021	
Mode of	IEC60601-1:2005	Continuous
operation	+A1:2012+A2:2020	
	EN 60601-1:2006	
	A1:2013+A2:2021	
Degree of	IEC 60529:1989+A1:1999	IP54 (provided by enclosures)
protection	EN 60529:1991+A1:2000	
General	93/42/EEC as amended by	Medical device Directive (class IIb)
	2007/47/EC	
	ISO 13485:2016	Quality systems - Medical devices -
	EN ISO 13485:2016	Requirements for regulating purposes
	ISO 14971:2019	Application of risk management to Medical
	EN ISO 14971:2019	devices
	IEC62304:2006+A1:2015	Medical device software - Software life-cycle
	EN 62304:2006/A1:2015	processes
	IEC60601-1-	Medical electrical equipment - Part 1-6:
	6:2010+A1:2013+A2:2020	General requirements for basic safety and
	EN60601-1-6:2010	essential performance - Collateral standard:
	+A1:2015+A2:2021	Usability
	IEC 62366-1:2015+A1:2020	Medical devices - Application of usability
	EN 62366-1:2015+A1:2020	engineering to Medical devices
	IEC60601-1-9:2007 +A1:2013	Medical electrical equipment - Part 1-9:
	EN 60601-1-9:2008 +A1:2013	General requirements for basic safety and
		essential performance - Collateral Standard:
		Requirements for environmentally conscious
	JEO 00004 4 44 0045 A4 0000	design
	IEC 60601-1-11:2015+A1:2020	Medical electrical equipment –
	EN 60601-1-11:2015+A1:2021	Part 1-11: General requirements for basic
		safety and essential performance –
		Collateral Standard: Requirements for
		medical electrical equipment and medical
		electrical systems used in the home
EMC	IEC 60604 4 0.0044 A4.0060	healthcare environment
EMC	IEC 60601-1-2:2014+A1:2020	Electromagnetic compatibility-requirements &
D-GL-90-4	EN 60601-1-2:2015+A1:2021	test
Defibrillator	IEC60601-2-4:2010+A1:2018	Medical electrical equipment - Part 2-4:
	EN60601-2-4:2011+A1:2019	Particular requirements for the basic safety
		and essential performance of cardiac
		defibrillators
Ambulance	EN 1789:2007 +A2:2014	Medical vehicles and their equipment
		 road ambulance

Item	Standard	Description
Package	ISTA (Procedure 2A, 2011)	Pre-Shipment test procedures (Package)
Battery	IEC60086-4:2014	Primary batteries - Part 4: Safety of lithium
		batteries
Marking	ISO 15223-1:2016	Medical devices. Symbols to be used with
	EN ISO 15223-1:2016	medical device labels, labelling and
		information to be supplied. General
		requirements
Labeling	EN 1041:2008	Information supplied by the manufacturer with
		Medical devices
	2021/2226/EU	Electronic instructions for use of medical
		devices (eIFU)
Hazardous	2015/863/EU	Restriction of the use of Hazardous
Substance		Substances in electrical and electronic
		equipment III (RoHS III)
	EC/1907/2006	Registration, Evaluation, Authorization and
		Restriction of Chemicals (REACH)
Disposal	2012/19/EU	Waste electrical and electronic equipment
		directive (WEEE)

Manufacturer's Declaration



For best product performance and measurement accuracy, use only accessories supplied or recommended by Mediana. Use accessories according to the manufacturer's directions for use and your facility's standards. The use of accessories, transducers, and cables other than those specified may result in increased emission and/or decreased immunity of the AED.

The AED is suitable for use in the specified electromagnetic environment. The customer and/or user of the AED should assure that it is used in an electromagnetic environment as described below:

Table 5. Electromagnetic Emissions (IEC60601-1-2)

Emission Test	Compliance	Electromagnetic Environment
RF emission	Group 1	The AED must emit electromagnetic energy
CISPR 11		in order to perform its intended function.
		Nearby electronic equipment may be
		affected.
RF emissions	Class B	The AED is suitable for use in all
CISPR 11		establishments.

Table 6. Electromagnetic Immunity (IEC60601-1-2)

Immunity Test	IEC60601-1-2 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	Floor should be wood, concrete, or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30 %.			
Power frequency (50 Hz) magnetic field IEC 61000-4-8	3 A/m	80 A/m	It may be necessary to position the AED further from the sources of power frequency magnetic fields or to install magnetic shielding. The power frequency magnetic field should be measured in the intended installation location to assure that it is sufficiently low.			
Note: UT is the AC	mains voltage prior to	Note: UT is the AC mains voltage prior to application of the test level.				

Table 7. Electromagnetic Immunity (IEC60601-1-2) (continued)

Immunity	IEC60601	Compliance	Electromagnetic
Test	test level	Level	environment guidance
			environment specified below. The sed in such an environment.
Radiated RF IEC 61000-4-3	10 V/m 80 MHz to 2.7 GHz (80 % AM at 1 kHz) (According to IEC60601-1- 2:2014) 10V/m, 20 V/m 80 MHz to 2.5 GHz (80 % AM at 1 kHz) (According to IEC60601-2- 4:2010)	10 V/m 20V/m	Potable and mobile RF communications equipment should be used no closer to any part of the AED including cables, than the recommended separation distance calculated from the equation appropriate to the frequency of the transmitter. Recommend separation distance d = 0.4 √P 80 MHz to 800 MHz d = 0.7 √P 800 MHz to 2.5 GHz d = 0.2 √P 80 MHz to 800 MHz d = 0.4 √P 800 MHz to 2.5 GHz 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 metres (m). Field strengths from fixed RF transmitters as deter-mined by an electromagnetic site survey, a should be less than the compliance level in each frequency range. Interference may occur in the vicinity of equipment marked with the following symbol:

Note: At 80 MHz and 800 MHz, the higher frequency range applies. Note: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Recommended Separation Distances between Portable and Mobile RF Communications Equipment and the AED. (IEC60601-1-2)

Note: According to the revised IEC60601-1-2:2014, there is no distinction between Life-Supporting device and not Life-Supporting device; all device shall ensure basic safety and essential performance.

^a Field strengths from fixed transmitters, such as base stations for radio (cellular/ cordless) telephones and land mobile 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 AED is used exceeds the applicable RF compliance level above, the AED should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as re-orienting or relocating the AED.

^b Over the frequency range 80 MHz to 2.5 GHz, field strengths should be less than 10 V/m

Table 8. Recommended Separation Distances

Recommended separation distance between Portable and mobile RF communications equipment and the AED

The AED is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the AED can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the AED as recommended below, according to the maximum output power of the communications equipment.

Rated Maximum Output Power of	Separation distance according to frequency of transmitter in meter		
Transmitter in watt	80 MHz to 800 MHz d = 1.2 √P	800 MHz to 2.5GHz d = 2.3 √P	
0.01	0.12	0.23	
0.1	0.38	0.73	
1	1.2	2.3	
10	3.8	7.3	
100	12	23	

For transmitters rated at a maximum output power not listed above, the recommended separation distance d in metres (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: At 80MHz and 800MHz, the separation distance for the higher frequency range applies Note: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects, and people.

Table 9. Immunity to proximity fields from RF wireless communications equipment (IEC 60601-1-2)

(IEC 60601-1-2)						
Test frequency (MHz)	Band ^a (MHz)	Service ^a	Modulation b	Maximum power (W)	Distance (m)	Immunity test level (V/m)
385	380 - 390	TETRA 400	Pulse Modulation: 18 Hz	1.8	1	27
450	430 - 470	GMRS 460, FRS 460	FM + 5Hz deviation 1 kHz sine	2	1	28
710 745 780	704 - 787	LTE Band 13, 17	Pulse Modulation: 217 Hz	0.2	1	9
810 870 930	800 - 960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	Pulse Modulation: 18 Hz	2	1	28
1720 1845 1970	1700 - 1990	GSM 1800; CDMA 1900; GSM 1900; DECT; LTE Band 1, 3, 4, 25; UMTS	Pulse Modulation: 217 Hz	2	1	28
2450	2400 - 2570	Bluetooth, WLAN, 802.11 b/g/n, RFID 2450, LTE Band 7	Pulse Modulation: 217 Hz	2	1	28
5240 5500 5785	5100 - 5800	WLAN 802.11 a/n	Pulse Modulation: 217 Hz	0.2	1	9

Note: If necessary to achieve the IMMUNITY TEST LEVEL, the distance between the transmitting antenna and the ME EQUIPMENT or ME SYSTEM may be reduced to 1 m. The

Table 10. Cables (IEC60601-1-2)

Cables and Sensors	Maximum Length	Complies with
Pads cable	1.8 m	-RF emissions, CISPR 11, Class B/ Group 1 -Electrostatic discharge (ESD), IEC 61000-4-2 -Radiated RF, IEC 61000-4-3 -Power frequency Magnetic field, IEC 61000-4-8

¹ m test distance is permitted by IEC 61000-4-3.

a For some services, only the uplink frequencies are included.
b The carrier shall be modulated using a 50 % duty cycle square wave signal.
c As an alternative to FM modulation, 50 % pulse modulation at 18 Hz may be used because while it does not represent actual modulation, it would be worst case.