HeartSine® samaritan® PAD
SAM 350P Semi-Automatic Defibrillator
SAM 360P Fully Automatic Defibrillator
SAM 450P Semi-Automatic Defibrillator
Use of This Manual

It is important that you read this manual carefully before using your samaritan® PAD. This manual is presented in support of any training you may have received. If you have any questions, contact your Authorized Distributor or HeartSine® Technologies directly.

CAUTION:
U.S. Federal law restricts this device to sale by or on the order of a licensed practitioner.

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Indications for Use

The HeartSine samaritan PAD SAM 350P (SAM 350P), HeartSine samaritan PAD SAM 360P (SAM 360P) and HeartSine samaritan PAD SAM 450P (SAM 450P) all have the identical indications for use. Each is indicated for use on victims of cardiac arrest who are exhibiting the following signs:

- Unconscious
- Not breathing
- Without circulation (without a pulse)

The devices are intended for use by personnel who have been trained in their operation. Users should have received training in basic life support/AED, advanced life support or a physician-authorized emergency medical response training program.

The devices are indicated for use on patients greater than 8 years old or over 55 lbs/25 kg when used with the adult Pad-Pak™ (Pad-Pak-01 or Pad-Pak-07). They are indicated for use on children between 1 and 8 years of age or up to 55 lbs/25 kg when used with the Pediatric-Pak™ (Pad-Pak-02).

Contraindications for Use

If the patient is responsive or conscious, do not use the samaritan PAD to provide treatment.

Caution

U.S. Federal law restricts this device to sale by or on the order of a physician.
WARNINGS

Fully Automatic Defibrillator (SAM 360P)
The SAM 360P is a fully automatic defibrillator. When required, it will deliver a shock to the patient WITHOUT user intervention.

CPR Rate Advisor Function (SAM 450P)
The CPR Rate Advisor function is intended for use on adult patients only. If a Pediatric-Pak is used, the CPR Rate Advisor function is disabled. In this case, the rescuer is prompted to begin CPR in time with the metronome but receives no CPR Rate Advisor feedback.

PRECAUTIONS

Correct Placement of Electrode Pads
Proper placement of the samaritan PAD electrode pads is critical. You must strictly observe the instructions shown on pages 19-22 and on the device. Wrong placement or the presence of air, hair, surgical dressings or medicine patches between the pads and the skin could reduce defibrillation effectiveness. Slightly red skin after shock therapy is normal.

Do Not Use Electrode Pads if Pouch is Not Sealed
The Pad-Pak and Pediatric-Pak are single-use items which must be replaced after each use or if the pouch that seals the electrode pads has been broken or compromised in any way. If you suspect that the Pad-Pak or Pediatric-Pak is damaged, replace it immediately.

Susceptibility to Electromagnetic Interference
To safeguard against interference, operate the samaritan PAD at least 6 feet/2 meters away from all radio frequency devices. Alternatively, switch off the equipment causing the electromagnetic interference.

Temperature Range for Operation
The samaritan PAD, with its battery and electrodes, is designed to operate in the temperature range of 32°F to 122°F/0°C to 50°C. Use of the device outside of this range may cause the device to malfunction.

Ingress Protection
The samaritan PAD has an IP56 rating against dust and sprays of water. However, the IP56 rating does not cover the immersion of any part of the samaritan PAD in water or any type of fluid. Contact with fluids may seriously damage the device or cause fire or a shock hazard.

Prolonging Battery Life
Do not turn on the device unnecessarily as this may reduce the standby life of the device.

Operation outside the range of 32°F to 122°F/0°C to 50°C may decrease the shelf-life of the Pad-Pak.

Operator Training
The samaritan PAD is intended for use by personnel who have been trained in its operation. Users should have received training in basic life support/AED, advanced life support, or a physician-authorized emergency medical response training program.

Use of Accessories
The samaritan PAD is a self-contained device. Do not use any unauthorized accessories with the device as the samaritan PAD may malfunction if non-approved accessories are used.

Regular Maintenance
Check the device periodically. See Service and Maintenance on page 27.

Correct Disposal of the Device
Dispose of the device in accordance with your national or local regulations, or contact your Authorized Distributor for assistance. Please follow the steps provided in After Using the samaritan PAD on page 25.

Compliance with Local Regulations
Check with the relevant local government health department for information about any requirements associated with ownership and use of a defibrillator in the region where it is to be used.

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Overview

Sudden Cardiac Arrest

Sudden cardiac arrest (SCA) is a condition in which the heart suddenly stops pumping blood effectively due to a malfunction of the heart’s electrical system. Often victims of SCA have no prior warning signs or symptoms. SCA also can occur in people with previously diagnosed heart conditions. Survival from SCA depends on immediate and effective cardiopulmonary resuscitation (CPR).

The use of an external defibrillator within the first few minutes of a collapse can greatly improve a patient’s chance of survival. Heart attack and SCA are not the same, though sometimes a heart attack can lead to an SCA. If you are experiencing symptoms of a heart attack (chest pain, pressure, shortness of breath, tight feeling in the chest or elsewhere in the body), immediately seek medical attention.

Sinus Rhythm and Ventricular Fibrillation

The normal heart rhythm, known as sinus rhythm, creates electrical activity resulting in coordinated contraction of the heart muscle. This generates normal blood flow around the body.

Ventricular fibrillation (V-fib or VF) is a condition in which there is uncoordinated contraction of the heart muscle, making it quiver rather than contract properly. Ventricular fibrillation is the most commonly identified arrhythmia in SCA patients. In victims of SCA it is possible to re-establish normal sinus rhythm by means of an electric shock across the heart. This treatment is called defibrillation.

Ventricular Tachycardia

Ventricular tachycardia (VT) is a type of tachycardia (rapid heartbeat) that arises from improper electrical activity of the heart. VT starts in the bottom chambers of the heart, called the ventricles. Although there are many different types of VT, this arrhythmia can be potentially life-threatening if the patient presents with no pulse and is unresponsive. If not treated with immediate defibrillation VT may lead to other arrhythmias.

Treatment by AED

It is a common misconception that CPR alone and calling emergency services is enough. CPR is a temporary measure that maintains blood flow and oxygen to the brain. CPR alone will not return a heart to a normal rhythm during VF or VT. The key to survival is defibrillation – and the sooner the better.

Defibrillation is a common treatment for life-threatening arrhythmias, mainly ventricular fibrillation. Defibrillation consists of delivering an electrical shock to the heart with a device called a defibrillator. This restores normal heart muscle contractions and allows normal sinus rhythm to be restored by the body’s natural pacemaker in the heart.

The samaritan PAD uses the HeartSine samaritan ECG arrhythmia analysis algorithm. This algorithm will evaluate the patient’s ECG to ascertain if a therapeutic shock is appropriate. If a shock is required, the samaritan PAD will charge and advise the user to press the shock button (SAM 350P/450P) or will automatically deliver a shock (SAM 360P). If no shock is advised, the device will pause to allow the user to deliver CPR.

It is important to note that cardiac defibrillators, like the HeartSine samaritan PAD, will not administer a shock unless a lifesaving shock is required.
This manual provides instructions for the following models of the HeartSine samaritan PAD:

- samaritan PAD 350P (SAM 350P)
- samaritan PAD 360P (SAM 360P)
- samaritan PAD 450P (SAM 450P)

### About the samaritan PAD

The samaritan PAD family of AEDs is designed to quickly deliver a defibrillation shock to victims of sudden cardiac arrest (SCA). Each samaritan PAD is designed to operate in accordance with the current joint American Heart Association (AHA) and European Resuscitation Council (ERC) guidelines on Cardiopulmonary Resuscitation (CPR) and Emergency Cardiovascular Care (ECC).

While all of the samaritan PAD models are very similar in use, there are distinct differences between the models as shown in Table 1 below.

#### Shock delivery

- **SAM 350P**: Semi-Automatic
- **SAM 360P**: Fully Automatic
- **SAM 450P**: Semi-Automatic

#### Four-year electrode and battery life

- **SAM 350P**: √
- **SAM 360P**: √
- **SAM 450P**: √

#### Audible and visual indicators

- **SAM 350P**: √
- **SAM 360P**: √
- **SAM 450P**: √

#### CPR coaching with metronome

- **SAM 350P**: √
- **SAM 360P**: √
- **SAM 450P**: √

#### CPR Rate Advisor

- **SAM 350P**: √
- **SAM 360P**: √
- **SAM 450P**: √

#### Pediatric use-compatible (with Pediatric Pad-Pak)

- **SAM 350P**: √
- **SAM 360P**: √
- **SAM 450P**: √

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### SAM 350P

The SAM 350P is a semi-automatic defibrillator, the SAM 360P is a fully automatic defibrillator, and the SAM 450P is a semi-automatic defibrillator with integrated CPR Rate Advisor™.

**WARNING:** The SAM 360P is a fully automatic defibrillator. When required, it will deliver a shock to the patient WITHOUT user intervention.

### CPR Metronome

When the samaritan PAD instructs you to perform CPR, you will hear an audible beep and see the Safe to Touch indicator flash at a rate compliant with the current joint American Heart Association (AHA) and European Resuscitation Council (ERC) guidelines on Cardiopulmonary Resuscitation (CPR) and Emergency Cardiovascular Care (ECC).

### CPR Rate Advisor

When providing CPR treatment to a victim of sudden cardiac arrest, it is vital the chest compressions are of a good quality. If the quality of the CPR provided is good, the chances of successfully resuscitating a patient are greatly increased.

Research has demonstrated that non-professional responders regularly provide ineffective CPR due to inexperience.

The SAM 450P with CPR Rate Advisor provides feedback to the rescuers on the rate of the CPR they are providing to the victim. The SAM 450P uses impedance cardiogram measurements to analyze the speed of compressions and provide the user with instructions to push faster or push slower or to continue to provide compressions at a good speed according to the AHA resuscitation guidelines. The SAM 450P uses both audible and visual feedback to give the responder instruction on CPR rate. Refer to Technical Data in Appendix C on page C-7.

**WARNING:** The CPR Rate Advisor function is intended for use on adult patients only. If a Pediatric-Pak is used, the CPR function is disabled.

In this case, the rescuer is prompted to begin CPR in time with the metronome but receives no CPR Rate Advisor feedback.

### Recommended Training

SCA is a condition requiring immediate emergency medical intervention. Due to the nature of the condition, this intervention can be performed before seeking the advice of a physician.

The samaritan PAD is intended for use by personnel who have been trained in its operation. Users should have received training in basic life support/AED, advanced life support, or a physician-authorized emergency medical response training program. HeartSine Technologies also recommends that this training be kept up to date by regular refresher courses as and when recommended by your training provider.

If potential users of the samaritan PAD are not trained in these techniques, contact your Authorized Distributor or HeartSine Technologies directly. Either can arrange for training to be provided. Alternatively contact your local government health department for information on certified training organizations in your area.

### Safety and Effectiveness Data

Please refer to Appendix E for the potential risks associated with the use of an AED and a summary of SAM 350P, SAM 360P and SAM 450P safety and effectiveness data.
SAM 450P Layout

Status Indicator
The SAM 450P is ready for use when this indicator is flashing green.

On/Off button
Press this button to turn on or turn off the device.

Safe to Touch Icon/Action Arrows
You may touch the patient when the action arrows around this icon are flashing.

Speaker
Listen for the metronome and verbal prompts.

Attach Pads Icon/Action Arrows
Attach the electrode pads to the patient’s bare chest as indicated when the action arrows are flashing.

Data Port
Plug the custom USB cable into this port to download event data from the AED. (See Figure 8, page 24.)

Shock Button
Press this button to deliver a therapeutic shock.

Adult and Pediatric Symbols
Indicates that the SAM 450P is compatible with both the Pad-Pak and Pediatric-Pak.

CPR Rate Advisor Icon
Provides visual feedback about the rate of chest compressions during CPR.

Do Not Touch Icon/Action Arrows
Do not touch the patient when the action arrows above this icon are flashing. The SAM 450P may be analyzing the patient’s heart rhythm or about to charge, in preparation to deliver a shock.

On/Off button
Press this button to turn on or turn off the device.

Pad-Pak
Contains the battery and electrode pads.

Green Tab
Pull this tab to release the electrodes.

Attach Pads Icon/Action Arrows
Attach the electrode pads to the patient’s bare chest as indicated when the action arrows are flashing.

Unpacking
Verify that the contents include the Samaritan PAD, carry case, Pad-Pak, User Manual, Warranty Statement and Warranty Card.

Pad-Pak
A Pad-Pak is a single-use removable cartridge that includes the battery and electrode pads in a single unit. The Pad-Pak is available in two versions:

1. Pad-Pak (gray color shown in Figure 1) for use on patients weighing over 55 lbs/25 kg, or equivalent to a child of approximately eight years of age or older.

2. The optional Pediatric-Pak (pink color shown in Figure 2) for use on smaller children (from 1 to 8 years old and weighing under 55 lbs/25 kg).

WARNING: Do not delay treatment trying to determine the patient's exact age and weight.

1 The Pad-Pak also is available in a TSO-certified version for use on aircraft.
Putting the samaritan PAD into Service

Follow these steps to place your samaritan PAD into service:

1. Check the expiration date (year-month-day) on the rear of the Pad-Pak (see Figure 3). If the expiration date has passed, do not use and immediately replace the expired Pad-Pak.

2. Unpack the Pad-Pak and retain the packaging in case you need to return the Pad-Pak to HeartSine Technologies.

3. Place the samaritan PAD face up on a flat surface and slide the Pad-Pak into the samaritan PAD (see Figure 4) until you hear the “double click” to indicate that the tabs on the right and left sides of the Pad-Pak are fully engaged.

4. Verify that the green Status indicator (see the layout for your model on pages 10-12) is blinking to indicate the initial self-test routine has been performed and the device is ready for use.

5. Press the On/Off Button to turn on the samaritan PAD. Listen for, but do not follow, the voice prompts to ensure that no warning messages are played.

6. Press the On/Off Button to turn off the samaritan PAD. Verify that the Status Indicator is flashing green. If you have not heard a warning message and the Status Indicator continues to flash green, the device is ready for use.

7. Place the samaritan PAD in its supplied soft carry case. Store the samaritan PAD where it will be seen and heard in an unobstructed, secure location in a clean, dry environment. Be sure to store the device according to the environmental specifications (see Technical Data in Appendix C on page C-1).

PRECAUTION: Do NOT pull the green tab on the Pad-Pak at this time. If you have pulled the tab and opened the electrode drawer, you may need to replace your Pad-Pak.

Only turn on the samaritan PAD ONCE. If you turn it on and off repeatedly, you will deplete the batteries prematurely and may need to replace the Pad-Pak.

8. Register online, or complete the Warranty Card and return it to your Authorized Distributor or HeartSine Technologies directly (see Tracking Requirements on page 26).

9. Create a service schedule (see Service and Maintenance on page 27).

PRECAUTION: HeartSine Technologies recommends that you store a spare Pad-Pak with your samaritan PAD in the rear section of the soft carry case.

Preparation Checklist

Following is a checklist of the steps required to set up your samaritan PAD:

- Step 1. Check the Pad-Pak expiration date.
- Step 2. Install the Pad-Pak and check for a green status indicator.
- Step 3. Turn on the samaritan PAD to check operation.
- Step 4. Turn off the samaritan PAD.
- Step 5. Store the samaritan PAD in a clean, dry environment at 32°F to 122°F/0°C to 50°C.
- Step 6. Register your samaritan PAD.
- Step 7. Create a service schedule. (See Service and Maintenance on page 27)
Using the samaritan PAD

Follow these steps to use your AED, which will provide you with step-by-step voice prompts. For a full list of voice prompts for your device see Voice Prompts in Appendix D.

**PRECAUTION:** Once a non-shockable rhythm is detected, the samaritan PAD will end its ready to shock condition if it had previously decided to shock.

1. If necessary, move the patient to a safe location, or remove any source of danger.

**PRECAUTION:** You must use the samaritan PAD at least 6 feet/2 meters from all radio frequency devices, or switch off any equipment causing electromagnetic interference.

2. If the patient is non-responsive, shake the patient by the shoulders while speaking loudly. If the patient becomes responsive, do not use the AED.

3. Check that the patient’s airway is not blocked, using a head-chin tilt if necessary.

4. Call for medical assistance.

5. Retrieve the AED, asking others nearby to do so.

6. While waiting for the AED, begin CPR, pushing hard and fast at a rate of between 100 and 120 compressions per minute (cpm) and a depth of 5 to 6 cm. If you feel able to give rescue breaths perform 30 compressions followed by two rescue breaths.

7. Press the On/Off Button to turn on the AED.

8. Defibrillation therapy is tailored depending on whether a Pad-Pak or Pediatric-Pak is installed. If the patient is under 55 lbs/25 kg or 8 years of age, remove the Pad-Pak, insert a Pediatric-Pak and press the On/Off button again (see Pediatric-Pak on page 21). If a Pediatric-Pak is not available, you may use the Pad-Pak.

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Using the samaritan PAD:  

9. Remove clothing from patient’s chest to expose bare skin, removing any metal (bras or jewelry) where possible from the pad placement area. 

10. Dry the patient’s chest if wet or clammy, and if a lot of chest hair is present, shave the patient’s chest where the electrodes will be placed.

11. Pull the green tab to remove the electrode pad pouch from the AED.

12. Tear open the pouch to remove the electrode pads.

13. Peel the liner from each electrode pad and apply each electrode pad firmly to the patient’s bare chest. For a patient over 8 years of age or weighing over 55 lbs/25 kg, place one electrode pad horizontally on the right chest, and the other vertically on the left rib cage. For a patient under 8 years of age or weighing less than 55 lbs/25 kg, you can place one electrode pad on the center of the chest and the other on the center of the back. Refer to pages 21-22 for detailed instructions for electrode pad placement.

14. If you again hear the prompt to apply pads firmly to patient’s bare chest, check that the:  
   - Pads are placed correctly as per pad placement shown.
   - Pads are not touching and at least 2.5 cm apart.
   - Entire surface of each pad is adhered to bare skin. If the chest is hairy, shave the chest; if the chest is wet, dry the chest.
   - Ensure the Pad-Pak has not expired, and is correctly inserted into the device.

15. When prompted, ensure that you are not touching the patient.
16. When advised that a shockable rhythm is detected, stand clear of patient as directed. When advised to do so, press the orange shock button (SAM 350P/SAM 450P) to deliver a shock, or if using a SAM 360P, the AED will automatically deliver the shock after a verbal 3, 2, 1 countdown.

17. When advised that a shockable rhythm is not detected, begin CPR. To do so, place overlapping hands in the middle of the patient’s chest and, with straight arms, press firmly and quickly in time with the metronome. Continue to perform CPR until the AED begins to analyze the patient’s heart rhythm again.

When using the SAM 450P, follow the CPR Rate Advisor voice prompts. Refer to CPR Rate Advisor on page C-7 for more information.

18. Repeat the process from step 15 until emergency services arrive.

19. When emergency services arrive, press the On/Off button to turn off the AED and remove the electrode pads.

Pediatric-Pak

Treating Small Children and Infants
The Pediatric-Pak is intended to provide therapy for pediatric (child) victims of SCA between the ages of 1 and 8 years old or weighing less than 55 lbs/25 kg who are:

- Unconscious
- Not breathing
- Without circulation (without a pulse)

WARNING: The Pediatric-Pak contains a magnetic component (surface strength 6500 gauss). Avoid storage next to magnetically-sensitive storage media.

WARNING: Not for use on patients under one year old. For use with children up to the age of 8 years or up to 55 lbs/25 kg. DO NOT DELAY THERAPY IF YOU ARE UNSURE OF THE EXACT AGE OR WEIGHT.

Electrode Placement
For pediatric patients there are two options for electrode placement: anterior-posterior and anterior-lateral.

ANTERIOR-POSTERIOR PLACEMENT
If a child’s chest is small it may be necessary to place one electrode pad in the centre of the child’s BARE chest (anterior), and the other electrode pad in the center of the ribcage on the child’s BARE back (posterior) as shown in Figure 5.
Pedicrian-Pak continued

ANTERIOR-LATERAL PLACEMENT
If a child’s chest is large enough to permit a 1 in/2.5 cm gap between the electrode pads, OR if trauma does not allow for placement on the back, the pads can be placed according to the adult anterior-lateral placement. Place one electrode pad on the child’s BARE upper right chest above nipple and one electrode pad on child’s BARE lower left ribcage below nipple as shown in Figure 6.

Figure 6. Anterior-Lateral Placement

WARNING: Electrode pads must be at least 1 in/2.5 cm apart and should never touch one another.

After Using the samaritan PAD

Cleaning the samaritan PAD
1. Remove the electrode pads from the patient and stick the pads together face to face. The electrodes may be contaminated with human bodily tissue, fluid or blood so dispose of the electrodes separately as infectious waste material.
2. The Pad-Pak is a single-use item that contains lithium batteries. Replace the Pad-Pak after each use. With the samaritan PAD placed face up on a flat surface, squeeze the two tabs on the sides of the Pad-Pak and pull to remove it from the samaritan PAD. The Pad-Pak will slide forward (see Figure 7).
3. Check the samaritan PAD for dirt or contamination. If necessary, clean the device using a soft cloth dampened by one of the following:
   • Soapy water
   • Isopropyl alcohol (70% solution)

PRECAUTION: Do not immerse any part of the samaritan PAD in water or any type of fluid. Contact with fluids may seriously damage the device or cause a fire or a shock hazard.

PRECAUTION: Do not clean the samaritan PAD with abrasive materials, cleaners or solvents.

4. Check the samaritan PAD for damage. If the device is damaged, replace it immediately.
5. Install a new Pad-Pak. Before installing the Pad-Pak, check the expiration date (see Set-up on page 14). After installation, confirm that the Status Indicator is blinking green.
6. Report the use of the samaritan PAD to HeartSine Technologies or your Authorized Distributor. (See back cover for contact details.)
After using the samaritan PAD

**Download and Submitting Event Information**

The optional HeartSine Saver EVO™ software can be downloaded at no charge from:


This software lets you manage the events in which your samaritan PAD was used. You can provide this data to a patient’s doctor, and/or use it to obtain a Pad-Pak if you have a qualifying event. In addition to Saver EVO, the optional USB data cable is required to download event data. Contact your Authorized Distributor or HeartSine Technologies directly to obtain the data cable or with questions about downloading and using Saver EVO.

1. Connect the USB data cable to the Data Port on the samaritan PAD (see Figure 8).

![Figure 8. USB Data Port](image)

2. Connect the USB connector on the data cable to a PC.

3. Install and launch the HeartSine Saver EVO software.

4. Follow the instructions provided in the Saver EVO manual to save or erase the event data on your samaritan PAD.

5. Upload the Saver EVO file on the HeartSine Technologies site.

For further information on managing the event data on your samaritan PAD, contact your Authorized Distributor or HeartSine Technologies directly.

**Disposal**

The Pad-Pak and Pediatric-Pak contain lithium batteries and cannot be disposed of in normal waste. Dispose of each at an appropriate recycling facility according to your local requirements. Alternatively return the Pad-Pak or Pediatric-Pak to your Authorized Distributor for disposal or replacement.
Tracking

Tracking Requirements
Medical device regulations require HeartSine Technologies to track the location of each Samaritan PAD AED, Pad-Pak, and Pediatric-Pak sold. Therefore, it is important that you register your device, either using our on-line registration tool at:

https://secure.heartsine.com/UserRegistration.html

Or by completing the Samaritan PAD Warranty Card and returning it to your Authorized Distributor or HeartSine Technologies directly. As an alternative to the card and on-line registration tool, you may send an email to:

support@heartsine.com

The email should contain the following information:
- Name
- Address
- Device Serial Number

If there is a change in the information you have provided to us, such as a change of address or ownership of your Samaritan PAD, provide the updated information to us via email or the online registration tool.

When you register your AED, we will contact you with any important notifications about the Samaritan PAD, such as software updates or field safety corrective actions.

Service and Maintenance

HeartSine Technologies recommends users perform regular maintenance checks, which include the following:

WEEKLY
- Check the Status Indicator. The Samaritan PAD performs a self-test routine at midnight GMT every Sunday. During this self-test the status light blinks red but returns to green upon successful completion of the self-test routine. If the Status Indicator is not flashing green every 5 to 10 seconds or if the status indicator is flashing red or you hear continuous beeping, a problem has been detected. (See Figures 9–11, and Troubleshooting in Appendix B on page B-1.)

MONTHLY
- If the device shows any signs of physical damage, contact your Authorized Distributor or HeartSine Technologies directly.
- Check the expiration date of the Pad-Pak (see Set-up on page 14 for the location of the date). If the date has expired, or is near expiration, immediately replace the Pad-Pak or contact your Authorized Distributor for a replacement.
- If you hear a warning message when you turn on your Samaritan PAD or if, for any reason, you suspect that your Samaritan PAD is not working properly, consult Troubleshooting in Appendix B.

Testing with Simulators and Manikins
HeartSine devices cannot be tested using industry-standard simulators and manikins. Therefore, to test the Samaritan PAD with a simulator or manikin, contact HeartSine Technologies or your Authorized Distributor for assistance.
APPENDIX A Symbols

Symbols Used in This Manual

![WARNING](https://example.com/warning-icon.png)
**WARNING:** Risk of death or serious injury

![PRECAUTION](https://example.com/precaution-icon.png)
**PRECAUTION:** Risk of injury

Symbols Used on the samaritan PAD

- **On/Off**
- **IP56** Ingress protection classified as IP56 according to EN 60529
- **Consult operating instructions**
- **Single use item; do not re-use**
- **Defibrillation protected, Type BF connection**
- **Do not incinerate or expose to high heat or open flame**
- **Does not contain natural rubber latex**
- **Non-sterile**
- **A-Recyclable**

**Non-rechargeable battery**

**Do not short circuit battery**

**Do not crush battery**

**Temperature limitation as indicated**

**Expiration date for Pad-Pak; yyyy-mm-dd**

**Dispose of in accordance with country requirements**

**Automated External Defibrillator**

With respect to electrical shock, fire and mechanical hazards only in accordance with:
- **ANSI/AAMI ES60601-1:2005**
- **CSA C22.2 NO. 60601-1:2008**
- **IEC60601-2-4:2010**

**Follow instructions for use**

**Serial number:** For example “yyB01234567” where yy = year of manufacture
# APPENDIX B Troubleshooting

## Flashing Red Status Indicator/Continual Beeping, or No Status Indicator Light is Lit

Check the expiration date on your Pad-Pak (see Set-up on page 14). If the expiration date has passed, immediately replace the Pad-Pak. If the expiration date has not passed, press the On/Off Button on the face to turn on the samaritan PAD and listen for the voice prompt “Call for medical assistance”. Then press the On/Off Button again to turn off the device. If either of these actions do not correct the problem, contact your Authorized Distributor or HeartSine Technologies immediately.

## “Low Battery” Warning

While this message does not indicate a fault, you should replace the battery as soon as possible. The first time you hear the message “Warning low battery,” the device will continue to function properly. However, it may have fewer than 10 shocks left so prepare the spare Pad-Pak for use and be prepared to swap it quickly. Order a new Pad-Pak as soon as possible.

## “Memory Full” Prompt

This message does not indicate a fault. The memory is full and can no longer record ECG data or events. However, the device can still analyze and deliver a shock if required. Contact HeartSine Technologies Technical Support for guidance on how to clear the memory.

## Three Rapid Beeps When Device Is Turned Off or After Weekly Self-Test Has Been Performed

Your device has sensed that the ambient temperature is outside the specified operating range. Return your device to the specified operating conditions of 32°F to 122°F/0°C to 50°C, in which your device, with its battery and electrodes is designed to operate, and verify that the beeping has stopped.

## Red Status Indicator and Beeping While Device is On

⚠️ **Warning:** There is insufficient battery capacity to deliver a shock. Immediately replace the Pad-Pak or seek an alternative defibrillator. If a spare Pad-Pak or alternative defibrillator is not available, the device will continue to analyze the patient’s heart rhythm and advise when CPR is needed, but it will not be able to deliver a shock.

## “Device service required” Warning

⚠️ **Warning:** If you hear this message during use, seek an alternative defibrillator immediately. Do not attempt to service the device as no modification of this equipment is possible. Contact HeartSine Technologies or your Authorized Distributor immediately.

## “You have pressed the off button” Warning

You have pressed the On/Off button while the AED is being used to treat a patient. If you are sure you want to turn off the AED, quickly press On/Off again.

## “Disarming” Prompt

This message does not indicate a fault; rather it means that the AED has converted to a decision to not shock after it has initially decided to shock. This occurs when your AED has initially determined that the patient’s rhythm is shockable (such as VF) and upon confirming the decision (before proceeding with a shock), the rhythm changed or interference (due to CPR) prevents the confirmation. Continue to follow the device prompts.
APPENDIX B Troubleshooting continued

Obtaining Support
If you have completed the troubleshooting steps and find the device is still not working correctly, contact your Authorized Distributor or HeartSine Technologies Technical Support at:
support@HeartSine.com

Warranty Exclusion
HeartSine Technologies or its Authorized Distributors are not obliged to replace or repair under warranty if one or more of the following conditions apply:

• Device has been opened.
• Unauthorized modifications have been made.
• Device has not been used in accordance with the instructions provided in this manual.
• Serial number has been removed, defaced, altered or, by any other means, made unreadable.
• Device has been used or stored outside its indicated temperature range.
• The Pad-Pak or Pediatric-Pak is not returned in its original packaging.
• Device has been tested using unapproved methods or inappropriate equipment (see Warnings and Precautions on pages 3-5).

APPENDIX C Technical Data

Physical Specifications (with Pad-Pak installed)
Size: 8.0 in x 7.25 in x 1.9 in/20 cm x 18.4 cm x 4.8 cm
Weight: 2.4 lbs/1.1 kg

Environmental Specifications
Operating temperature: 32°F to 122°F/0°C to 50°C
Standby temperature: 32°F to 122°F/0°C to 50°C
Transport temperature: 14°F to 122°F/-10°C to 50°C for up to two days. If the device has been stored below 32°F/0°C, it should be returned to an ambient temperature of between 32°F to 122°F/0°C to 50°C for at least 24 hours before use.

Relative humidity: 5% to 95% (non-condensing)
Enclosure: IEC/EN 60529 IP56
Altitude: 0 to 15,000 feet/0 to 4,575 meters
Shock: MIL STD 810F Method 516.5, Procedure 1 (40G’s)
Vibration: MIL STD 810F Method 514.5+, Procedure 1
Category 4 Truck Transportation – US Highways
Category 7 Aircraft – Jet 737 & General Aviation

Physical Specifications
Size: 8.0 in x 7.25 in x 1.9 in/20 cm x 18.4 cm x 4.8 cm
Weight: 2.4 lbs/1.1 kg

Environmental Specifications
Operating temperature: 32°F to 122°F/0°C to 50°C
Standby temperature: 32°F to 122°F/0°C to 50°C
Transport temperature: 14°F to 122°F/-10°C to 50°C for up to two days. If the device has been stored below 32°F/0°C, it should be returned to an ambient temperature of between 32°F to 122°F/0°C to 50°C for at least 24 hours before use.

Relative humidity: 5% to 95% (non-condensing)
Enclosure: IEC/EN 60529 IP56
Altitude: 0 to 15,000 feet/0 to 4,575 meters
Shock: MIL STD 810F Method 516.5, Procedure 1 (40G’s)
Vibration: MIL STD 810F Method 514.5+, Procedure 1
Category 4 Truck Transportation – US Highways
Category 7 Aircraft – Jet 737 & General Aviation
### Pad-Pak and Pediatric-Pak Specifications

**Weight:** 0.44 lbs/0.2 kg  
**Battery type:** Disposable single-use combined battery and defibrillation electrode cartridge (lithium manganese dioxide (LiMnO2) 18V)  
**Battery capacity (new):** >60 shocks at 200J or 6 hours of continuous monitoring  
**Battery capacity (4 years):** >10 shocks at 200J  
**Electrode type:** Single-use pre-attached combined ECG sensor/defibrillation pad  
**Electrode placement:**  
- **Adult:** Anterior-lateral  
- **Pediatric:** Anterior-posterior or anterior-lateral  
**Electrode active area:** 15 in² / 100 cm²  
**Electrode cable length:** 3.3 feet / 1 m  
**Shelf life/Standby life:** See the expiration date on the Pad-Pak/Pediatric-Pak  

### Aircraft Safety Test (TSO-Certified Pad-Pak):  
RCAA DO-227 (ETSO-C142a)

### Patient Analysis System  
Method: Evaluates the patient’s ECG, signal quality, electrode contact integrity and patient impedance to determine if defibrillation is required  
**Sensitivity/Specificity:** Meets IEC/EN 60601-2-4 (Refer to page C-6 for sensitivity/specificity data.)

### User Interface  
**Visual prompts:** Adult and Pediatric Symbols, Do Not Touch Icon/Action Arrows, Safe to Touch Icon/Action Arrows, Status Indicator, Attach Pads Icon/Action Arrows, CPR Rate Advisor Indicator (SAM 450P only)  
**Audible prompts:** Extensive voice prompts guide the user through the operation sequence (see Voice Prompts in Appendix D)  
**Languages:** samaritan PAD is available in English and Spanish. Contact your HeartSine Authorized Distributor.  
**Controls:** On/Off Button (all models), Shock Button (SAM 350P and 450P only) and Green Tab

### Defibrillator Performance  
**Charging time:** Typically 150J in < 8 seconds, 200J in < 12 seconds  
**Time to shock delivery following CPR:**  
- **SAM 350P:** Typically 8 seconds  
- **SAM 360P:** Typically 19 seconds  
- **SAM 450P:** Typically 12 seconds  
**Impedance range:** 20 Ω to 230 Ω

### Therapeutic Shock  
**Waveform:** SCOPE™ (Self Compensating Output Pulse Envelope) optimized biphasic escalating waveform compensates energy, slope and envelope for patient impedance  
**Energy:** Pre-configured factory settings for escalating energy are based on the current AHA/ERC guidelines  
- **Pad-Pak:**  
  - Shock 1: 150J; Shock 2: 150J; Shock 3: 200J  
- **Pediatric-Pak:**  
  - Shock 1: 50J; Shock 2: 50J; Shock 3: 50J

### Event Recording  
**Type:** Internal memory  
**Memory:** 90 minutes of ECG (full disclosure) and event/incident recording  
**Review:** Custom USB data cable (optional) directly connected to a PC with Saver EVO Windows-based data review software

### Electromagnetic Compatibility/Battery Safety  
**EMC:** IEC/EN 60601-1-2  
**Radiated emissions:** IEC/EN 55011  
**Electrostatic discharge:** IEC/EN 61000-4-2 (8 kV)  
**RF Immunity:** IEC/EN 61000-4-3 80 MHz – 2.5 GHz, (10 V/m)  
**Magnetic field immunity:** IEC/EN 61000-4-8 (3 A/m)  
**Aircraft:**  
- RTCA/DO-160G, Section 21 (Category M)  
- RTCA DO-227 (ETSO-C142a)
SCOPE Biphasic Waveform

The samaritan PAD delivers a Self-Compensating Output Pulse Envelope (SCOPE) biphasic waveform which automatically optimizes the waveform pulse envelope (amplitude, slope, and duration) for a wide range of patient impedances, from 20 ohms to 230 ohms. The delivered waveform to the patient is an optimized, impedance-compensated, biphasic, truncated exponential waveform that incorporates an escalating energy protocol of 150 joules, 150 joules, and 200 joules. The duration of each phase is automatically adjusted to compensate for varying patient impedances. The first phase (T₁) duration is always equivalent to the second phase (T₃) duration. The interphase pause (T₂) is always a constant 0.4 ms for all patient impedances.

The specific SCOPE waveform characteristics for a 150 joules pulse are shown in Table 2. An example of waveform parameters for the Pediatric-Pak are shown in Table 3.

Table 2. Pad-Pak Waveform Specification

<table>
<thead>
<tr>
<th>Resistance (ohms)</th>
<th>Waveform Voltages (volts)</th>
<th>Waveform Duration (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>V₁</td>
<td>Tilt %</td>
<td>T₁</td>
</tr>
<tr>
<td>V₂</td>
<td>V₃</td>
<td>T₂</td>
</tr>
<tr>
<td>V₃</td>
<td></td>
<td>T₃</td>
</tr>
</tbody>
</table>

- 25 1640 63.1 3 3
- 50 1650 52.7 4.5 4.5
- 75 1660 51.4 6.5 6.5
- 100 1670 48.7 8 8
- 125 1670 50.4 10.5 10.5
- 150 1670 48.7 12 12
- 175 1670 48.7 14 14
- 200 1670 47.6 15.5 15.5
- 225 1680 46.7 17 17

NOTE: All values are nominal.

Table 3. Pediatric-Pak Waveform Specification

<table>
<thead>
<tr>
<th>Resistance (ohms)</th>
<th>Energy (joules)</th>
<th>Waveform Voltages (volts)</th>
<th>Waveform Duration (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>V₁</td>
<td>Tilt %</td>
<td>T₁</td>
<td></td>
</tr>
<tr>
<td>V₂</td>
<td>V₃</td>
<td>T₂</td>
<td></td>
</tr>
<tr>
<td>V₃</td>
<td>T₃</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- 25 47.5 51.4 3 3
- 50 51.3 671 4.5 4.5
- 75 52.1 751 6.5 6.5
- 100 51.8 813 10.5 10.5
- 125 52.4 858 14 14

NOTE: Table contains example Pediatric-Pak waveform parameters.

Motion Detection Algorithm (SAM 360P only)

The SAM 360P uses the samaritan PAD ICG analysis to detect chest compression artefact and other forms of motion in order to play a verbal warning to stop CPR or other motion.

If the algorithm detects motion or other significant interference, the SAM 360P will issue the voice prompt "Motion detected, do not touch the patient." This is intended to reduce the likelihood that the user is touching the patient prior to shock delivery.

NOTE: Motion Detection Algorithm performance may be reduced during low battery operation.
Arrhythmia Analysis Algorithm

The Samaritan PAD uses its ECG arrhythmia analysis algorithm to evaluate the patient’s ECG to determine if a therapeutic shock is appropriate. If a shock is required, the Samaritan PAD will charge and advise the user to stand clear and to press the shock button (SAM 350P and 450P) or automatically shock the patient after a verbal 3, 2, 1 countdown (SAM 360P). If no shock is advised, the device will pause to allow the user to deliver CPR.

The Samaritan PAD ECG arrhythmia analysis algorithm performance has been extensively evaluated by using several databases of real-life ECG traces. Included in this are the AHA database and the Massachusetts Institute of Technology (MIT) NST database. The Samaritan PAD ECG arrhythmia analysis algorithm’s sensitivity and specificity meet the requirements of IEC/EN 60601-2-4.

The Samaritan PAD ECG arrhythmia analysis algorithm performance is summarized in Table 4.

Table 4. Performance of the Samaritan PAD ECG Arrhythmia Analysis Algorithm

<table>
<thead>
<tr>
<th>Rhythm Class</th>
<th>Minimum Test Sample Size</th>
<th>Test Sample Size</th>
<th>Performance Goal</th>
<th>Observed Performance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shockable Rhythm: Coarse Ventricular Fibrillation</td>
<td>200</td>
<td>350</td>
<td>Sensitivity &gt; 90%</td>
<td>Met</td>
</tr>
<tr>
<td>Shockable Rhythm: Rapid Ventricular Tachycardia</td>
<td>50</td>
<td>53</td>
<td>Sensitivity &gt; 75% (AAMI DF39)</td>
<td>Met</td>
</tr>
<tr>
<td>Non-Shockable Rhythm: NSR*</td>
<td>100</td>
<td>165</td>
<td>Specificity &gt; 99% (exceeds AAMI DF39)</td>
<td>Met</td>
</tr>
<tr>
<td>Non-Shockable Rhythm: AF, SB, SVT, Heart Block, Idioventricular, PVCs*</td>
<td>30</td>
<td>153</td>
<td>Specificity &gt; 95% (from AAMI DF39)</td>
<td>Met</td>
</tr>
<tr>
<td>Non-Shockable Rhythm: Asystole</td>
<td>100</td>
<td>117</td>
<td>Specificity &gt; 95%</td>
<td>Met</td>
</tr>
<tr>
<td>Intermediate: Fine Ventricular Fibrillation</td>
<td>25</td>
<td>46</td>
<td>Report Only &gt;45% Sensitivity</td>
<td></td>
</tr>
<tr>
<td>Intermediate: Other Ventricular Tachycardia</td>
<td>25</td>
<td>29</td>
<td>Report Only &gt;65% Specificity</td>
<td></td>
</tr>
</tbody>
</table>

1. AAMI Association for Advancement of Medical Instrumentation; NSR, normal sinus rhythm; AF, atrial fibrillation/flutter; SB, sinus bradycardia; SVT, supraventricular tachycardia; PVCs, premature ventricular contractions.

CPR Rate Advisor

The Samaritan 450P utilizes the ICG (Impedance Cardiogram) capability to assess the rate of chest compressions being applied during cardiopulmonary resuscitation (CPR). Based on the measured rate, the Samaritan 450P provides verbal feedback to the user to “push faster”, “push slower”, or continue to provide “good speed” in accordance with the current AHA resuscitation guidelines (target CPR rate of at least 100 CPM).

The Samaritan 450P also uses the ICG to provide CPR Rate Advisor feedback in the form of a colored traffic light (green-amber-red) configuration LED array as shown in Table 5 below. The LED array is accompanied by text on the device’s top cover membrane to indicate when the operator’s compressions are ‘too slow’ or ‘too fast’.

Table 5. Visual guidance for CPR Rate Advisor

<table>
<thead>
<tr>
<th>Description</th>
<th>Audio message</th>
<th>LED sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>N/A</td>
<td>“Begin CPR”</td>
<td>[Red]</td>
</tr>
<tr>
<td>Compression rate is slower than 90 per minute</td>
<td>“Push faster”</td>
<td>[Green, Amber]</td>
</tr>
<tr>
<td>Compression rate is between 90 and 99 per minute</td>
<td>Metronome only</td>
<td>[Green]</td>
</tr>
<tr>
<td>Compression rate is between 100 and 120 per minute</td>
<td>“Good speed”</td>
<td>[Amber]</td>
</tr>
<tr>
<td>Compression rate is between 121 and 130 per minute</td>
<td>Metronome only</td>
<td>[Green]</td>
</tr>
<tr>
<td>Compression rate is faster than 130 per minute</td>
<td>“Push slower”</td>
<td>[Red]</td>
</tr>
</tbody>
</table>
Pediatric Restriction

Use of the CPR Rate Advisor function is restricted to adult patients only. Chest compression techniques differ for the different ages and sizes of pediatric patients (up to eight years old). For younger pediatric patients, rescuers should compress the lower half of the sternum but not compress over the xiphoid. For patients at the upper end of the pediatric range, adult-style compressions should be performed. CPR Rate Advisor is currently configured only to advise compressions at a rate suitable for adult patients (over eight years old weighing more than 55 lbs/25 kg).

Electrode placement also may differ in pediatric patients. Depending on the patient size, the electrodes may be placed anterior-posterior (front and back) or anterior-lateral (standard adult placement). Differing electrode positions may result in different ICG readings. Current technology does not support CPR Rate Advisor in determining which electrode placements are being used and therefore electrodes must be placed anterior-lateral for CPR Rate Advisor to function correctly.

For these reasons, CPR Rate Advisor is disabled when a Pediatric-Pak is used in the SAM 450P.

NOTE: The ECG readings used to determine if the patient requires a defibrillation shock are not affected by the electrode positions selected in pediatric patients.

WARNING: If a pediatric patient is treated with an adult Pad-Pak, ignore the CPR Rate Advisor feedback prompts provided. CPR Rate Advisor is currently only intended to provide feedback on adult patients.

Guidance and Manufacturer’s Declaration – Electromagnetic Emissions

The samaritan PAD is intended for use in the electromagnetic environment specified in Table 6. The customer or the user of the samaritan PAD should assure that it is used in such an environment.

Table 6. Electromagnetic Emissions

<table>
<thead>
<tr>
<th>Emissions Test</th>
<th>Compliance</th>
<th>Electromagnetic Environment – Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>RF emissions CISPR II</td>
<td>Group I</td>
<td>The SAM 350P, SAM 360P and SAM 450P use RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.</td>
</tr>
<tr>
<td>RF emissions CISPR II</td>
<td>Class B</td>
<td>The SAM 350P, SAM 360P and SAM 450P are suitable for use in all establishments, including domestic and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.</td>
</tr>
<tr>
<td>Harmonic emissions IEC/EN 61000-3-2</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Voltage fluctuations/flicker emissions IEC/EN 61000-3-3</td>
<td>Not applicable</td>
<td></td>
</tr>
</tbody>
</table>

Guidance and Manufacturer’s Declaration – Electromagnetic Immunity

The samaritan PAD is intended for use in the electromagnetic environment specified in Table 7. The customer or the user of the samaritan PAD should assure that it is used in such an environment.
### Table 7. Electromagnetic Immunity

<table>
<thead>
<tr>
<th>Immunity Test</th>
<th>IEC 60601 Test Level</th>
<th>Compliance Level</th>
<th>Electromagnetic Environment – Guidance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electrostatic discharge (ESD)</td>
<td>± 6 kV contact</td>
<td>± 6 kV contact</td>
<td>Floors should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.</td>
</tr>
<tr>
<td></td>
<td>± 8 kV air</td>
<td>± 8 kV air</td>
<td></td>
</tr>
<tr>
<td>Electrical fast transient/burst</td>
<td>±2kV for power supply lines</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td></td>
<td>±1 kV for input/output lines</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Surge</td>
<td>±1kV differential mode</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td></td>
<td>±2 kV common mode</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>Voltage dips, short interruptions and voltage variations on power supply input lines (IEC/EN 61000-4-11)</td>
<td>&lt;5 % U, (&gt;95 % dip in U) for 0.5 cycle</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td></td>
<td>40 % U, (60 % dip in U) for 5 cycles</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td></td>
<td>70 % U, (90 % dip in U) for 25 cycles</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td></td>
<td>&lt;5 % U, (&gt;95 % dip in U) for 5 sec</td>
<td>Not Applicable</td>
<td>Not Applicable</td>
</tr>
<tr>
<td>NOTE: U, is the a.c. mains voltage prior to application of the test level</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Power-frequency (50/60 Hz) magnetic field (IEC/EN 61000-4-8)</td>
<td>3 A/m</td>
<td>3A/m</td>
<td>Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.</td>
</tr>
<tr>
<td>Conducted RF (IEC/EN 61000-4-6)</td>
<td>3 V rms</td>
<td>Not applicable</td>
<td>Recommended separation distance</td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz outside ISM bands</td>
<td>Not applicable</td>
<td>Not applicable</td>
</tr>
<tr>
<td></td>
<td>10 V rms</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td></td>
<td>150 kHz to 80 MHz in ISM bands</td>
<td>Not applicable</td>
<td></td>
</tr>
<tr>
<td>Radiated RF (IEC/EN 61000-4-3)</td>
<td>10 V/m</td>
<td>10 V/m</td>
<td>Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, should be less than the compliance level in each frequency range.</td>
</tr>
<tr>
<td></td>
<td>80 MHz to 2.5 GHz</td>
<td>80 MHz to 2.5 GHz</td>
<td>Interference may occur in the vicinity of equipment marked with the following symbol:</td>
</tr>
</tbody>
</table>

NOTE:

- UT is the a.c. mains voltage prior to application of the test level.
- Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.
- Portable and mobile RF communications equipment should be used no closer to any part of the SAM 350P, SAM 360P or SAM 450P, including cables, than the recommended separation distance calculated from the equation applicable to the frequency of the transmitter.
- Field strengths from fixed RF transmitters, as determined by an electromagnetic site survey, 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 1: At 80 MHz and 800 MHz, the higher frequency range applies.

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

a. The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

b. The compliance levels in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz are intended to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas. For this reason, an additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance for transmitters in these frequency ranges.

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

Recommended Separation Distances between Portable and Mobile RF Communication Equipment and the samaritan PAD

The samaritan PAD is intended for use in an electromagnetic environment in which radiated RF disturbances are controlled. The customer or the user of the samaritan PAD can help prevent electromagnetic interference by maintaining a minimum distance between portable and mobile RF communications equipment (transmitters) and the samaritan PAD as recommended in Table 8, according to the maximum output power of the communications equipment.

```
<table>
<thead>
<tr>
<th>Rated Maximum Output Power of Transmitter W</th>
<th>Separation Distance According to Frequency of Transmitter m</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>150 kHz to 80 MHz Outside ISM Bands</td>
</tr>
<tr>
<td>0.01</td>
<td>Not applicable</td>
</tr>
<tr>
<td>0.1</td>
<td>Not applicable</td>
</tr>
<tr>
<td>1</td>
<td>Not applicable</td>
</tr>
<tr>
<td>10</td>
<td>Not applicable</td>
</tr>
<tr>
<td>100</td>
<td>Not applicable</td>
</tr>
</tbody>
</table>

For transmitters rated at a maximum output power not listed above, the recommended separation distance (d) in meters (m) can be estimated using the equation applicable to the frequency of the transmitter, where P is the maximum output power rating of the transmitter in watts (W) according to the transmitter manufacturer.

APPENDIX C Technical Data continued
APPENDIX D Voice Prompts

Following are the voice prompts used by the samaritan PAD devices. Models using specific voice prompts are indicated. Read the voice prompts in advance of use to be familiar with the types of instructions given.

<table>
<thead>
<tr>
<th>PROMPT</th>
<th>SAM 350P</th>
<th>SAM 360P</th>
<th>SAM 450P</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Call for medical assistance”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Remove clothing from patient’s chest to expose bare skin”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Pull green tab to remove pads”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Peel pads from liner”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Apply pads to patient’s bare chest as shown in picture”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Press pads firmly to patient’s bare skin”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Assessing heart rhythm; do not touch the patient”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Analyzing; do not touch the patient”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
<tr>
<td>“Motion detected”</td>
<td>✔️</td>
<td>✔️</td>
<td>✔️</td>
</tr>
</tbody>
</table>

NOTE 1: At 80 MHz and 800 MHz, the separation distance for the higher frequency range applies.

NOTE 2: The ISM (industrial, scientific and medical) bands between 150 kHz and 80 MHz are 6.765 MHz to 6.795 MHz; 13.553 MHz to 13.567 MHz; 26.957 MHz to 27.283 MHz; and 40.66 MHz to 40.70 MHz.

NOTE 3: An additional factor of 10/3 has been incorporated into the formulae used in calculating the recommended separation distance for transmitters in the ISM frequency bands between 150 kHz and 80 MHz and in the frequency range 80 MHz to 2.5 GHz to decrease the likelihood that mobile/portable communications equipment could cause interference if it is inadvertently brought into patient areas.

NOTE 4: These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.
### Voice Prompts continued

<table>
<thead>
<tr>
<th>PROMPT</th>
<th>SAM 350P</th>
<th>SAM 360P</th>
<th>SAM 450P</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CPR Rate Advisor</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Push faster”*</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Push slower”*</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Good speed”*</td>
<td></td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td><strong>If a Shock is Not Required</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>“No shock advised”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Begin CPR”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“It is safe to touch the patient”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Place overlapping hands in middle of chest”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Press directly down on the chest in time with metronome”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Remain calm”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td><strong>If a Shock is Required</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>“Stand clear of patient; shock advised”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Stand clear of patient; press the orange shock button now”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Stand clear of patient; shock will be delivered in 3, 2, 1”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Shock delivered”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Begin CPR”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“It is safe to touch the patient”</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Place overlapping hands in middle of chest”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Press directly down on the chest in time with metronome”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>“Remain calm”*</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

*Voice prompts not provided when Pediatric-Pak is installed.

### Potential Adverse Effects

The potential adverse effects (e.g., complications) associated with the use of an automated external defibrillator include, but are not limited to, the following:

- Failure to identify shockable arrhythmia.
- Failure to deliver a defibrillation shock in the presence of VF or pulseless VT, which may result in death or permanent injury.
- Inappropriate energy which could cause failed defibrillation or post-shock dysfunction.
- Myocardial damage.
- Fire hazard in the presence of high oxygen concentration or flammable anesthetic agents.
- Incorrectly shocking a pulse-sustaining rhythm and inducing VF or cardiac arrest.
- Bystander shock from patient contact during defibrillation shock.
- Interaction with pacemakers.
- Skin burns around the electrode placement area.
- Allergic dermatitis due to sensitivity to materials used in electrode construction.
- Minor skin rash.

### Overall Safety Summary

It is difficult to determine the percentage of the intended population that could expect to experience a harmful event without a prospective, randomized clinical trial. In lieu of such a trial, the risks of the device are based on nonclinical laboratory and animal studies as well as data collected in published literature.

The results from the preclinical laboratory testing performed on the HeartSine SAM 350P, SAM 360P and SAM 450P devices demonstrated appropriate electrical safety, electromagnetic compatibility, biocompatibility, mechanical integrity, and overall performance.

Six animal studies were conducted to demonstrate device safety. The first study involved 15 pigs and demonstrated first shock success of the SCOPE waveform used in the SAM 350P, SAM 360P and SAM 450P of 99%.
Two animal studies were conducted to determine whether the SCOPE waveform induces refibrillation or increases defibrillation thresholds (DFTs) if the pulse duration exceeds 20 ms. The first of these studies included 6 pigs and the delivery of 208 shocks, including 68 shocks with pulse durations greater than or equal to 20 ms; the second pulse width study included 5 pigs and 96 delivered shocks, of which 91 shocks had pulse durations greater than or equal to 20 ms. There were no incidents of refibrillation in the 5 minutes immediately following a successful first shock in either study. In addition, successful defibrillation demonstrated that the SCOPE waveform at longer pulse durations does not increase DFTs.

The fourth animal study was conducted to assess the impact of administering repeated shocks on a pig model wearing an underwire bra. In this study, the bra’s underwire was intentionally exposed by removal of bra fabric and the electrode pad was placed so that the electrode gel on the electrode’s lower surface was in direct contact with the bra’s metal wire, to maximize the potential of arcing or other adverse events. A total of 126 shocks were administered at the device’s maximum energy of 200 J, with 100% first shock success and no evidence of arcing, redirection of current away from the subject, scorching or burning to the animal or fabric, or any other skin damage to the animal observed post-resuscitation.

The fifth animal study demonstrated the accuracy of the SAM 360P audio feedback (“Motion detected”, “Do not touch the patient” and “Stand clear”) by determining if CPR motion/interference which was intentionally created during the ECG rhythm analysis period was detected by the SAM 360P Motion Detection Algorithm. This 12-pig study demonstrated a sensitivity for the “motion detected” audio feedback of 97.9% and a specificity of 100%.

The sixth animal study demonstrated the accuracy of the SAM 450P CPR Rate Feedback algorithm. This 12-pig study demonstrated that the proportion of correct audio and visual feedback provided by the SAM 450P was 95.2% in comparison to the actual CPR compression rate as demonstrated in video recordings.

The animal studies collectively demonstrate: the safety of the devices’ energy protocol; that pulse durations greater than 20 ms do not induce refibrillation or increase DFTs; that placement of the electrodes in proximity with a bra does not pose a risk; and that the SAM 360P motion/interference detection algorithm and SAM 450P CPR Rate Feedback features do not adversely affect CPR quality or shock delivery.

Overall Effectiveness Summary

In addition to the bench and animal studies described previously, bench testing demonstrates that the arrhythmia analysis algorithm meets AHA recommendations for sensitivity and specificity for detecting shockable and non-shockable arrhythmias. A published clinical study on the SCOPE waveform, postmarket clinical experience involving 805 events, and usability studies conducted on each device demonstrate the effectiveness of the SAM 350P, SAM 360P and SAM 450P devices.

Published Clinical Data

The study by Walsh et al. evaluated the SCOPE defibrillation waveform used in the SAM 350P, SAM 360P and SAM 450P and was published in the American Journal of Cardiology in 2004. This study compared two (2) impedance compensated biphasic waveforms:

- HeartSine’s samaritan (SAM) (100-150-200 J energy protocol at the time based on the AHA guidelines then in effect) using the same SCOPE defibrillation waveform present in the SAM 350P, SAM 360P and SAM 450P.
- Philips Medical Systems Heartstream XL (150-150-150 J protocol) (HSXL)

The primary endpoint was discontinuation of ventricular arrhythmia. Success was defined as the discontinuation of ventricular arrhythmia for greater than 5 seconds. Patients were excluded from participation if they weighed less than 36 kg, had cardiac arrest due to trauma or had current “do not resuscitate” instructions.

As reported in the publication, 78 consecutive patients were studied: 40 HSXL (19 men) and 38 SAM (28 men). Mean age was 69 ± 11 years for HSXL patients and 65 ± 14 years for SAM patients (p = NS). Cardiac arrest out-of-hospital occurred in 13 of 40 HSXL patients (33%) and in 26 of 38 SAM patients (68%) (p = 0.003). Mean response time from arrest to physician arrival was 1.4 ± 1.3 minutes for in-hospital patients and 9 ± 6 minutes for out-of-hospital patients.

The rhythm when first recorded was VF in 20 of 40 HSXL patients (50%) and 16 of 38 SAM patients (42%), VT in 3 of 40 HSXL patients (8%) and 1 of 38 SAM patients (3%), and electromechanical dissociation or asystole in 16 of 40 HSXL patients (40%) and 20 of 38 SAM patients (53%) (all rhythms, p = NS). One (1) patient in each group had a palpable pulse when first attended by the physician. Drugs given during cardiac arrest were similar in the two (2) groups. A total of 15 of 40 HSXL patients (38%) and 12 of 38 SAM patients (31%) had a pulse at the time of the first shock.
patients (32%) received amiodarone, whereas 29 of 40 HSXL patients (73%) and 34 of 38 SAM patients (89%) received epinephrine (p = NS).

VF episodes were 107 HSXL and 117 SAM. The energy selection protocol was adhered to in 95 of 107 HSXL (89%) and 79 of 117 SAM (68%) defibrillation episodes. Protocol violations for energy selection occurred when the attending physician misinterpreted a successful shock followed by early recurrence of arrhythmia (greater than 5 seconds) as unsuccessful. This resulted in a progression to the next stage of the energy selection protocol (i.e., a higher energy was therefore selected inappropriately). Less incorrect energy selection was seen with the HSXL due to the non-escalating nature of the protocol (150-150-150 J; the physician could only select 200 J at the fourth shock or beyond).

Excluding VF episodes when energy selection was not as per protocol, success after one (1) shock was seen for 64% of HSXL and 58% of SAM episodes (p = NS). Success occurred by shock two (2) in 78% of HSXL and 82% of SAM episodes and by shock three (3) in 83% of HSXL and 92% of SAM episodes.

An analysis of the difference of proportions in success by a certain shock was performed by the authors (see Table 9). The defibrillation success rate is acceptable because it is consistent with defibrillation success rates (greater than 85%) reported in the literature for randomized controlled clinical trials using other devices and waveforms4. These data were not powered to demonstrate differences in return of spontaneous circulation or survival.

This study was conducted on the HeartSine samaritan AED (cleared under 510(k) K023854) with the identical SCOPE waveform as used in the SAM 350P, SAM 360P and SAM 450P.

### Table 9: Summary of Successful Defibrillation Episodes with Both Devices

<table>
<thead>
<tr>
<th></th>
<th>HeartSine samaritan (100-150-200J)</th>
<th>Philips HeartstreamXL (150-150-150J)</th>
<th>Mean Difference</th>
<th>SD</th>
<th>Probability that samaritan Better than Heartstream</th>
</tr>
</thead>
<tbody>
<tr>
<td>Success by</td>
<td>Frequency</td>
<td>Proportion</td>
<td>Frequency</td>
<td>Proportion</td>
<td></td>
</tr>
<tr>
<td>First Shock</td>
<td>46</td>
<td>0.582</td>
<td>61</td>
<td>0.642</td>
<td>-0.0598</td>
</tr>
<tr>
<td>Second Shock</td>
<td>65</td>
<td>0.823</td>
<td>74</td>
<td>0.779</td>
<td>0.0438</td>
</tr>
<tr>
<td>Third Shock</td>
<td>73</td>
<td>0.924</td>
<td>79</td>
<td>0.832</td>
<td>0.0925</td>
</tr>
<tr>
<td>Total Episodes</td>
<td>79</td>
<td>95</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*p = 0.029

### Postmarket Clinical Data

In addition to the published clinical study data described above, postmarket clinical data were received and analyzed from 28 countries worldwide including USA, Singapore, Germany, Netherlands, Canada, Australia, United Kingdom, and Sweden which comprised approximately 85% of the total number of events.

Postmarket clinical reports for 805 events were received between January 2012 and December 2015. Of these, 550 (68.3%) events involved the SAM 300P device, 122 (15.2%) events involved the SAM 350P device, three (3) (0.4%) events involved the SAM 360P device, no (0%) events involved the SAM 450P device, and 130 (16.1%) events involved the SAM 500P device. The SAM 500P is not marketed in the United States but uses the identical defibrillation waveform, the identical arrhythmia detection algorithm and identical Pad-Paks. The SAM 300P is the precursor to the SAM 350P and also uses identical defibrillation waveform, arrhythmia detection algorithm and Pad-Paks.

Success was defined as discontinuation of ventricular fibrillation or ventricular tachycardia within 5 seconds of shock delivery. A total of 334 patients in the “All Cases” dataset initially presented with a shockable rhythm, of which 327 (97.9%) patients were in ventricular fibrillation and seven (7) (2.1%) were in ventricular tachycardia.
were in ventricular tachycardia. Of these 334 patients, a shock was delivered in 322 patients. Of these 322 events, 220 (68.3%) events involved the SAM 300P device, 37 (11.5%) events involved the SAM 350P device, 2 (0.6%) events involved the SAM 360P device, no (0%) events involved the SAM 450P device and 63 (19.6%) events involved the SAM 500P device.

Of the 322 first shocks delivered, 293 (91.0%) were successful, with 95% CI estimated to be (87.3%, 93.9%). This is consistent with defibrillation success rates (greater than 85%) reported in the literature for randomized controlled clinical trials using other devices and waveforms.4

Of the “Shock Delivered” dataset, a total of 187 (58.1%) patients were reported to have survived to hospital admission, 61 (18.9%) patients did not survive to hospital admission, and survival information was unavailable for 74 (23.0%) patients. Table 10 summarizes the relationship between event location and user training, response time, and percentage survival to hospital admission.

Table 10: Location of Events, Trained Users, Response Time and Survival

<table>
<thead>
<tr>
<th>Location of Events</th>
<th>N</th>
<th>Percentage of Total Number of Shock Delivered Events</th>
<th>Percentage of Trained Users</th>
<th>Mean (SD) Response Time (minutes)</th>
<th>Percentage Survival to Hospital Admission</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home</td>
<td>44</td>
<td>13.7</td>
<td>88.6</td>
<td>4.97 (2.56)</td>
<td>34.1</td>
</tr>
<tr>
<td>Medical Facility</td>
<td>28</td>
<td>8.7</td>
<td>93.3</td>
<td>4.07 (4.85)</td>
<td>67.9</td>
</tr>
<tr>
<td>Office</td>
<td>19</td>
<td>5.9</td>
<td>68.4</td>
<td>3.86 (2.97)</td>
<td>68.4</td>
</tr>
<tr>
<td>Public</td>
<td>110</td>
<td>34.2</td>
<td>81.8</td>
<td>3.82 (3.95)</td>
<td>69.1</td>
</tr>
<tr>
<td>School/University</td>
<td>4</td>
<td>1.2</td>
<td>100.0</td>
<td>4.00</td>
<td>75.0</td>
</tr>
<tr>
<td>Sports Facility</td>
<td>62</td>
<td>18.9</td>
<td>79.0</td>
<td>4.30 (5.40)</td>
<td>80.6</td>
</tr>
<tr>
<td>Unknown</td>
<td>57</td>
<td>17.7</td>
<td>30.3</td>
<td>5.59 (2.00)</td>
<td>11.8</td>
</tr>
<tr>
<td>Total</td>
<td>322</td>
<td>100.0</td>
<td>73.0</td>
<td>4.21 (4.11)</td>
<td>58.1</td>
</tr>
</tbody>
</table>

First shock success was found to be significantly associated with survival to hospital admission, with an Odds Ratio (OR) = 3.13, 95% CI = (1.30, 7.51), and p = 0.0107. The analysis was repeated when adjusting for age and gender, with consistent results (OR = 3.29, p = 0.0059). Age was found to be significantly associated with survival to admission in this analysis (OR = 0.98 for a -year increase in age, p = 0.0324).

Shock success and survival were similar among the HeartSine public access defibrillators studied in this analysis, which was anticipated since all the devices use the same defibrillation waveform, the same arrhythmia detection algorithm, and the same Pad-Pak electrode-battery packs. Table 11 summarizes shock success and survival by defibrillator model.

Table 11: Shock Success and Survival to Hospital Admission

<table>
<thead>
<tr>
<th>Device Type</th>
<th>Number of Patients with a Shockable Rhythm</th>
<th>Percentage First Shock Success (%)</th>
<th>Percentage Second Shock Success (%)</th>
<th>Percentage Third Shock Success (%)</th>
<th>Percentage Survival to Hospital Admission (“All Cases” Dataset) (%)</th>
<th>Percentage Survival to Hospital Admission of Those Who Received a Shock (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SAM 300P</td>
<td>225</td>
<td>91.3</td>
<td>89.2</td>
<td>79.4</td>
<td>26.2</td>
<td>55.0</td>
</tr>
<tr>
<td>SAM 350P</td>
<td>41</td>
<td>89.2</td>
<td>76.5</td>
<td>90.9</td>
<td>22.1</td>
<td>62.2</td>
</tr>
<tr>
<td>SAM 360P</td>
<td>2</td>
<td>100.0</td>
<td>100.0</td>
<td>100.0</td>
<td>33.3</td>
<td>50.0</td>
</tr>
<tr>
<td>SAM 450P</td>
<td>0</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>SAM 500P</td>
<td>66</td>
<td>92.1</td>
<td>82.8</td>
<td>76.9</td>
<td>37.7</td>
<td>66.7</td>
</tr>
<tr>
<td>Total</td>
<td>334</td>
<td>91.3</td>
<td>86.4</td>
<td>81.4</td>
<td>27.5</td>
<td>58.1</td>
</tr>
</tbody>
</table>
The primary adverse event was failure to deliver a shock when presented with a shockable rhythm. A total of 12 of the 334 patients initially presented with a shockable rhythm but no shock was delivered. Based on a review of the ECG records, in only 1 of the 12 cases was algorithm performance determined to be inappropriate. In addition, 12 events were associated with audio prompts indicating the user repeatedly removed the electrodes throughout the event. No other adverse events were observed in this postmarket experience.

In summary, the postmarket data collected provides information on the real-world performance of the arrhythmia detection algorithm, waveform effectiveness and the overall usability of HeartSine’s public access defibrillators. First shock success and survival to admission were comparable in this study to rates reported in published literature. Finally, algorithm performance for combined VF/VT had a sensitivity of 98.8% in this analysis.

REFERENCES